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(54) Title: IMMUNOCONJUGATES COMPRISING SINGLE-CHAIN VARIABLE REGION FRAGMENTS OF ANTI-CD-19 ANTI-BODIES

(57) Abstract

Disclosed are polynucleotides encoding single chain variable region fragments of a monoclonal antibody to CD19 and methods for preparing the same. Also disclosed are single chain variable region polypeptides, methods for preparing the same, point modified polypeptides, and dimers derived therefrom. An additional aspect of the invention discloses immunoconjugates formed between a polypeptide of the invention and cytotoxic agents, as well as methods for their preparation, as well as use in the treatment of cancer.

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IMMUNOCONJUGATES COMPRISING SINGLE-CHAIN VARIABLE REGION FRAGMENTS OF ANTI-CD-19 ANTIBODIES

POSSIBLE GOVERNMENT OWNERSHIP RIGHTS

The research leading to the information disclosed herein was supported by the National Institutes of Health (NIH) under Grant No. CA49721. As a consequence, the government of the United States of America may possess certain rights to the invention disclosed herein.

BACKGROUND OF THE INVENTION

Immunoconjugates

Antibodies directed against cell surface molecules defined by cluster differentiation (CD) antigens represent a unique opportunity for the development of therapeutic reagents. Certain CD antigen expression is highly restricted to specific lineage lymphohematopoietic cells and, over the past several years, antibodies directed against lymphoid-specific CD antigens have been used to develop treatments that were effective either *in vitro* or in animal models (Ghetie *et al.*, 1988; Uckun *et al.*, 1986; Myers *et al.*, 1991; Jansen *et al.*, 1992). However, due to their large size, intact antibodies and antibody-toxin conjugates have several disadvantages that limit their efficiency. They are restricted in their ability to migrate from

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the vascular system, are heterogeneous as immunoconjugates (which can result in linkage of several toxin molecules to one immunoglobulin molecule), and their production is expensive and very labor intensive. See, for example, U.S. Patent No. 4,831,117 to Uckun and U.S. Patent No. 4,671,958 to Rodwell, *et al.*, the teachings of which are herein incorporated specifically by reference.

The limited efficacy of many unmodified monoclonal antibodies has led to an alternative approach, the use of these agents as carriers of cytotoxic substances. An array of toxins of bacterial and plant origin have been coupled to monoclonal antibodies for production of immunotoxins (Schlom; Pastan et al., 1986). The strategy is to select from nature a toxic protein and then to modify the toxin so that it will no longer indiscriminately bind and kill normal cells but will instead kill only the cells expressing the antigen identified by the monoclonal antibody. The majority of toxins targeted to cell surfaces by immunoconjugates act in the cytoplasm, where they inhibit protein synthesis. After binding to cell surface antigens, immunotoxins are taken up by endocytosis and delivered to endosomes. Fragments of some toxins (for example, diphtheria toxin) are then translocated across the membrane of this organelle. Other immunotoxins (for example, ricin) are routed further to the trans-Golgi network, where a minority undergo translocation to the cytoplasm. Unfortunately, most are routed to lysosomes, where they are degraded. In the cytoplasm, the toxins used clinically act either to adenosine diphosphate (ADP)-ribosylate elongation factor 2 (for example, Pseudomonas exotoxin (PE)) or to inactivate the 60S ribosomal subunit so that it has a decreased capacity to bind elongation factor 2 (for example, ricin). Less than ten toxin molecules in the cytoplasm are sufficient to kill the cell; however, more must bind to the cell surface to compensate for the inefficiencies in internalization and translocation.

Although immunotoxins are simple in concept, the first-generation immunotoxins were relatively ineffective. Several requirements must be fulfilled for an immunotoxin to be effective (Pastan *et al.*, 1986). In

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particular: (i) the immunoconjugate should be specific and should not react with normal tissues. Binding to tissues that do not express antigen can be reduced by removal of the nonspecific natural cell-binding subunits or domains of the toxin. Furthermore, because plant glycoprotein toxins contain mannose oligosaccharides that bind to cells of the reticuleondothelial system and, in some cases, also contain fucose residues that are recognized by the receptors on hepatocytes, deglycosylation of plant toxins may be required to avoid rapid clearance and potential cytotoxic effects on these cells. (ii) The linkage of the toxin to the antibody should not impair the capacity of the antibody to bind antigen. (iii) The immunotoxin must be internalized into endosomic vesicles. Thus, toxins directed by monoclonal antibodies to surface receptors that are normally internalized may be more active than those directed toward noninternalizing cell surface molecules. (iv) The active component of the toxin must translocate into the cytoplasm. These various goals can be in conflict; thus, the removal of the B chain of ricin reduces nonspecific binding but also reduces the capacity of the residual A-chain monoclonal antibody conjugate to translocate across the endosomic vesicle membrane. (v) For in vivo therapy, the linkage must be sufficiently stable to remain intact while the immunotoxin passes through the tissues of the patient to its cellular site of action. The first generation of heterobifunctional crosslinkers used to bind the toxin to the monoclonal antibody generated disulfide bonds that were unstable in vivo. This problem was solved in part by the synthesis of more stable cross-linkers, which used phenyl or methyl groups, or both, adjacent to the disulfide bond to restrict access to the bond.

The activity of an immunotoxin is initially assessed by measuring its ability to kill cells with target antigens on their surfaces. Because toxins act within the cells, receptors and other surface proteins that naturally enter cells by endocytosis usually make good targets for immunotoxins, but surface proteins that are fixed on the cell surface do not. However, if several antibodies recognizing different epitopes on the same cell surface

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protein are available, it is useful to test them all, because some, perhaps by producing a conformational change in the target protein's structure, may induce its internalization or direct its intracellular routing to an appropriate location for toxin translocation (May et al., 1991; Press et al., 1988). Also, it is possible to induce internalization of a target surface protein if the immunotoxin contains a form of PE or ricin in which the binding of the toxin moiety to its receptor, although weakened by chemical modification, still occurs and promotes internalization since toxin receptors are efficiently internalized (Willingham et al., 1987; Lambert et al., 1991; Colombatti et al., 1986).

Several immunotoxins have been developed and approved for human trials. Two different kinds of trials have been conducted. The first involves the ex vivo addition of immunotoxins to harvested bone marrow to eliminate contaminating tumor cells before reinfusion in patients undergoing autologous bone marrow transplantation. A variety of antibodies, linked to ricin or ricin A chain, including anti-CD5 and anti-CD7, have been used for this purpose (Uckun et al., 1990b). The second kind of trial involves the parenteral administration of immunotoxins, either regionally (such as the peritoneal cavity) or systematically, to patients with cancer. These have been primarily Phase 1 and 2 trials in patients in which conventional treatments have failed, and the patients have a large tumor burden. So far, the antibodies used for the preparation of immunotoxins to treat carcinomas or other solid tumors have been found to react with important normal human tissues (such as neural tissue and bone marrow) and produce dose-limiting toxicity without significant clinical responses (Weiner et al., 1989; Gould et al., 1989; Byers et al., 1989; Pai, in press).

Cell Differentiation Antigens

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The maturation of human BCPs into functional B lymphocytes represents a developmentally programmed multi-step process, which is accompanied by a cascade of somatic immunoglobulin gene rearrangements (Korsmeyer *et al.*, 1981), as well as a coordinated

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acquisition and loss of B-lineage differentiation antigens (Nadler). The characterization and classification of these antigens have been standardized during the first (Paris, France, 1982), second (Boston, MA, 1984), third (Oxford, UK, 1986), and fourth (Vienna, Austria, 1989) International Workshops on Human Leukocyte Differentiation Antigens, and a World Health Organization (WHO)-established CD (cluster of differentiation) nomenclature has been introduced for their identification (Nadler; Knapp *et al.*, 1989a; Clark *et al.*, 1989).

To date, more than 20 biochemically distinct differentiation antigens have been identified on B-lineage cells not including the surface immunoglobulins (sIg), major histocompatibility (MHC) antigens, or the receptor proteins for defined cytokines. Many of the B-lineage differentiation antigens represent functionally important surface receptors on developing B-lineage cells, and their expression is regulated by different external signals (Knapp et al., 1989a; Clark et al., 1989; Zola, 1987). While some (such as CD10, CD45, and CD73) represent membrane-associated enzymes, others (such as CD19, CD22, and B7) likely represent physiologically important cell surface bound ligands, which may play an important role in cell-to-cell interactions during B-cell development in a bone marrow microenvironment (Knapp et al., 1989a; Clark et al., 1989; Zola, 1987). The latter possibility is precedented by published evidence showing that many T-lineage differentiation antigens including CD2, CD4, CD8, and CD18/LFA-1 function as cell-surface bound ligands (CD2 for LFA-3, CD4 for class II MGC, CD8 for class I MHC, CD18/LFA-1 for I-CAM-1/gp80). The heterophilic recognition between such surface receptors may be important for cognate surface interactions between B-lineage cells and T cells or accessory cell populations in lymphohematopoietic tissues. Other B-lineage antigens (such as CD23 and CD40) might function as surface receptors for as yet undefined soluble cytokines (Clark et al., 1989).

CD19, CD22, and B7 antigens are members of the Ig supergene family (Knapp *et al.*, 1989b; Stamenkovic, 1988; Stamenkovic, 1990; Freeman *et al.*, 1989). CD21 has been identified as the C3d receptor as well

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as a receptor for Epstein-Barr virus (EBV) (Knapp et al., 1989b). The cytoplasmic domain of CD19 shows homology to proteins encoded by the int-1 oncogene and by EBV (Stamenkovic, 1988). CD19 has been proposed as a bridging molecule important for transduction of sIg-mediated signals in mature B cells (Pesando et al., 1989; Carter et al., 1990). CD19 as a signaltransducing subunit and CD21 as a ligand-binding subunit linking the B cell to the complement system have been reported to form a complex on the surface of B cells which may be involved in the sIg-dependent activation. However, the function of the CD19 molecule is not dependent on the presence of sIg or CD21 because CD19 ligation results in stimulation of phosphoinositide turnover (Uckun et al., 1989) and calcium mobilization in sIg-CD21-BCP populations and modulates their proliferative activity (Uckun et al., 1988; Ledbetter et al., 1988). CD22 displays a high degree of homology to the myelin-associated glycoprotein (MAG), a neuronal surface adhesion molecule mediating cell-to-cell interactions between B cells and monocytes (Stamenkovic, 1990. Furthermore, CD22 may also be important for transduction of sIgmediated signals (Pezzutto et al., 1988). Most recently, the natural ligand of B7 antigen has been identified as the CD28 T-cell activation antigen, which is another member of the Ig superfamily (Linsley et al., 1990). CD28-B7 mediated adhesion between activated B cells and T cells might be important for T-cell regulation of antigen-specific B-cell responses.

Monoclonal Antibodies and Fragments

Monoclonal antibodies have largely been applied clinically to the diagnosis and therapy of cancer and the modulation of the immune response to produce immunosuppression for treatment of autoimmune and graft versus host diseases (GVHD) and for prevention of allograft rejection. Human monoclonal antibodies have also been applied clinically against cytomegalovirus, *Varicella zoster* virus, and the various specific serotypes of *Pseudomonas aeruginosa*, *Escherichia coli*, and *Klebsiella pneumoniae*.

Antibodies or their fragments can also be genetically engineered to

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have more rapid clearance. This is desirable when a monoclonal antibody is conjugated to a radionuclide for use in radioimmunoscanning. For example, antigen-binding fragment (Fab), F(ab')₂, or single chain Fv fragments of monoclonal antibodies have survival half-lives of less than 5 hours. Rapid turnover can also be accomplished by the deletion of the CH2 domain as demonstrated for an antibody reactive with the disaloganglioside GD2 expressed on human tumors of neuroectodermal origin (Müeller *et al.*, 1990).

In an attempt to improve on the efficacy of anti-tumor cytotoxicity of antibodies and immunoconjugates, several laboratories have developed strategies for the expression of the light and heavy chain variable regions of antibodies in bacteria as single chain Fv (scFv) fragments (Pastan et al., 1991; Huston et al., 1988). In general, these molecules have been insoluble and need to be denatured and refolded before binding activity can be detected. One problem with production of antibody binding domains in this manner is that high affinity antibody binding cannot be successfully reconstituted in all instances. The parameters that govern the ability of an antibody to yield an scFv that can bind its target are unknown, thus necessitating the direct cloning and analysis of the candidate antibody gene segments.

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The CD19 antigen, which is found on mature B cells but not on plasma cells, has proven to be a very useful target for development of immunoconjugates because most lymphomas and B lineage leukemias express this differentiation marker (Uckun *et al.*, 1990a). Anti-CD19 immunoconjugates have relied on the chemical conjugation of the antibody and a modified catalytic toxin such as the A chain of ricin (Ghetie *et al.*, 1988) or pokeweed antiviral protein (Uckun *et al.*, 1986; Myers *et al.*, 1991). Prior to the development of the present invention, there have been no reports of the development of a successful scFv directed against the CD19 antigen.

The ability of immunotoxins to kill specific subsets of cells efficiently *in vitro* has led to their application in the deletion of particular

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cell types in suspensions of bone marrow cells (Thorpe et al., 1982; Seon, 1984; Vallera et al., 1982; Filipovich et al., 1984; Vallera et al., 1983; Muirhead et al., 1983; Krolick et al., 1982). The ultimate objective is to facilitate bone marrow transplantation in the human as an approach to treatment of cancer and diseases of the hematopoietic system. Autologous bone marrow transplantation is used as an adjunct to treatment for certain types of cancer which are highly susceptible to X-irradiation and or chemotherapy (Thomas, 1982; Raso, 1982). The approach is to obtain bone marrow from a patient in remission (preferably in the first remission) and to freeze it. If the patient subsequently relapses, the patient is then subjected to "supralethal" therapy with X-irradiation and or chemotherapy in order to eradicate the tumor. The patient is then rescued from death by infusion of his own bone marrow.

It would, of course, be highly desirable to purge such bone marrow of cancer cells by a cancer cell-reactive immunotoxin. The only requirement of such an immunotoxin is that it should not damage the stem cells which are needed to reconstitute the patient's hematopoietic system.

Immunoconjugates may be utilized for *ex vivo* purging of neoplastic cells from patient bone marrow grafts. These autologous grafts are reintroduced into leukemic patients after aggressive supra lethal chemotherapy and irradiation. The objective of all strategies is to deplete neoplastic cells while leaving unharmed the pluripotent hematopoietic stem cells which repopulate the patient's marrow after reinfusion. Intact immunoconjugates selectively eliminate antigen-positive targets without endangering engraftment and without causing intoxication.

Autologous marrow may be purged of residual leukemia cells without destroying hematopoietic stem cells by the use of immunoconjugates either *in vivo* or *ex vivo*. *Ex vivo* treatment with immunoconjugates has been shown to eliminate most T or B cells present in human marrow without damaging the ability of the marrow to reconstitute lethally irradiated recipients. While the efficiency of

immunoconjugates to kill "the last" leukemic cells still remains an issue the even greater efficiency of radiolabeled immunoconjugates should greatly increase the chances of successful treatment.

Radiolabeled Immunoconjugates

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It has been reported that an immunotoxin can specifically eliminate more than 99.99% of clonogenic leukemic T cells even in the presence of excess human bone marrow. The use of a radiolabeled immunotoxin should eliminate even more leukemic T cells, possibly at a rate of greater than 5 logs or 99.999%, indicating that the radiolabeled immunotoxin may be extremely useful for the *ex vivo* elimination of leukemic cells in autologous BMT.

Radiolabeled monoclonal antibodies have been developed as alternative immunoconjugates for delivery of a cytotoxic effector to target cells and for radioimaging (Schlom; Kozak *et al.*, 1985). These species possess potential to compensate for the observed shortcomings of immunotoxins. Toxin conjugates do not pass easily from the endosome to the cytosol. Furthermore, the toxins are immunogenic and thus provide only a short therapeutic window before the development of antibodies directed toward the toxin.

Radioimmunodetection with the use of radiolabeled monoclonal antibodies, most often with monoclonal antibodies to carcinoembryonic antigen, is widely used to complement other approaches for tumor detection. Although intact IgG antibodies are retained better by tumors and thus appear to be better for therapy, $F(ab')_2$ and Fab fragments are preferred for imaging because both targeting and blood clearance are most rapid, which reduces the background. Tumors as small as 0.5 cm, which are sometimes missed by other radiological methods, can be imaged with antibodies or antibody fragments labeled with suitable radionuclides.

One advantage in the use of radiolabeled monoclonal antibody conjugates for therapy is that with the appropriate choice of radionuclide, radiolabeled monoclonal antibodies can kill cells from a distance of several cell diameters and may therefore kill antigen-negative cells adjacent to

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antigen-expressing cells. Furthermore, the radiolabeled antibody need not be internalized to kill the tumor cell. Such techniques are exemplified in the teachings of U.S. Patent No. 4,831,122 to Buschbaum *et al.*, incorporated herein by reference.

In a radiolabeled monoclonal antibody, the radionuclide must be tightly linked to the antibody either directly or by a bifunctional chelate. For a monoclonal antibody-chelate complex to be effective, it must meet criteria in addition to those that are true for all monoclonal antibodies: (i) the chelating agent coupled to the monoclonal antibody should not compromise antibody specificity; (ii) the chelation and radiolabeling procedure should not alter the distribution and catabolism of the monoclonal antibody; and (iii) the bifunctional chelate should not permit elution and thus premature release of the radiolabeled metal *in vivo*. Failure to fulfill this last requirement has led to unacceptable toxicity and reduced efficacy. There are a number of suitable ∞ -, β -, and γ -emitting radionuclides. Isotopes emitting β particles, although superior to γ -emitting radionuclides, are not optimal because their low linear energy transfer released over a relatively long distance results in inefficient local killing of target cells coupled with toxicity to distant normal tissues.

Nevertheless, ß-emitting radionuclides such as ¹³¹I, ⁹⁰Y, ¹⁸⁸Re, and ⁶⁷Cu have been useful in immunotherapy. For example, hepatomabearing patients have been successfully treated with ¹³¹I-labeled antibodies to ferritin (Order, 1985). Furthermore, ⁹⁰Y-labeled antibodies to ferritin combined with autologous marrow transplantation resulted in complete remissions in four of eight patients with Hodgkin's disease (Order, 1985). ⁹⁰Y-labeled anti-Tac was effective in prolonging the survival of cardiac allografts and xenografts in a subhuman primate model (Kozak *et al.*, 1989). In a subsequent trial, ⁹⁰Y-labeled anti-Tac was evaluated for the treatment of patients with HTLV-I-associated, Tac-expressing ATL. At the doses used (5 and 10 miCi per patient), no toxicity was observed in five of six patients studied; modest granulocytopenia and thrombocytopenia were observed in one patient. Five of these six patients underwent a sustained

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partial or complete remission after 90Y-labeled anti-Tac therapy.

The target CD19 antigen, a 95 kDa B lineage restricted phosphoglycoprotein, is not expressed on life-maintaining non-hematopoietic tissues, normal hematopoietic progenitor cells, or most immature normal B-lineage lymphoid progenitor cells, but it is expressed by virtually 100% of B lineage ALLs.

SUMMARY OF THE INVENTION

In a first aspect, the present invention provides an isolated and purified polynucleotide encoding a single chain variable region polypeptide that binds to a CD19 antigen. Preferably, the isolated and purified polynucleotide of the invention encodes a polypeptide that has a molecular weight of approximately 28 kDa. More preferably, the polynucleotide of the invention encodes a polypeptide that binds to a CD19 antigen with a K_a of at least 1×10^9 M-1.

Also provided by the invention is a process for preparing an isolated and purified polynucleotide encoding a single chain variable region polypeptide that binds to a CD19 antigen comprising the steps of (a) isolating RNA from hybridomas producing monoclonal antibodies to CD19 antigen; (b) transcribing isolated RNA to cDNA; (c) amplifying separate cDNA sequences encoding a heavy and a light variable region of the monoclonal antibody by a polymerase chain reaction; (d) cloning of separate amplification products encoding the heavy and the light variable regions, respectively, into vector constructs; (e) cloning of DNA encoding the heavy and the light variable regions, wherein the separate sequences are joined through a linker nucleotide sequence, into a vector construct; and (f) digesting the clones with appropriate restriction endonucleases. Preferably, the process of the invention utilizes a linker sequence that is the polynucleotide sequence described by SEQ ID NO:7.

The present invention also contemplates an isolated and purified polynucleotide preparable by a process comprising the steps of (a) isolating RNA from hybridomas producing monoclonal antibodies to CD19 antigen;

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(b) transcribing isolated RNA to cDNA; (c) amplifying separate cDNA sequences encoding a heavy and a light variable region of the monoclonal antibody by a polymerase chain reaction; (d) cloning of separate amplification products encoding the heavy and the light variable regions, respectively, into vector constructs; (e) cloning of DNA encoding the heavy and the light variable regions, wherein the separate sequences are joined through a linker nucleotide sequence, into a vector construct; and (f) digesting the clones with appropriate restriction endonucleases.

Alternatively, the present invention provides an isolated and purified polynucleotide prepared by a process comprising the steps of (a) isolating RNA from hybridomas producing monoclonal antibodies to CD19 antigen; (b) transcribing isolated RNA to cDNA; (c) amplifying separate cDNA sequences encoding a heavy and a light variable region of the monoclonal antibody by a polymerase chain reaction; (d) cloning of separate amplification products encoding the heavy and the light variable regions, respectively, into vector constructs; (e) cloning of DNA encoding the heavy and the light variable regions, wherein the separate sequences are joined through a linker nucleotide sequence, into a vector construct; and (f) digesting the clones with appropriate restriction endonucleases. Preferably, the process of the invention utilizes a linker sequence that is the polynucleotide sequence described by SEQ ID NO:7.

Preferably, the isolated and purified polynucleotide of the claimed invention encodes a polypeptide comprising an amino acid residue sequence according to SEQ ID NO:20, 21 or 22. More preferably, the isolated and purified polynucleotide of the invention comprises a nucleotide sequence according to SEQ ID NO:23, 24 or 25.

In an alternative embodiment, the present invention provides an isolated and purified polynucleotide comprising a nucleotide base sequence that is identical or complimentary to a segment of at least 10 contiguous nucleotide bases of SEQ ID NO:23, 24 or 25, wherein the polynucleotide hybridizes to a polynucleotide that encodes a single chain variable region polypeptide that binds to a CD19 antigen. Preferably, the

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polynucleotide of this embodiment of the invention encodes a polypeptide that has a molecular weight of approximately 28 kDa. More preferably still, the encoded polypeptide binds to a CD19 antigen with a K_a of at least 1 x 10^9 M-1.

In another aspect, this embodiment of the invention provides an isolated and purified polynucleotide comprising a nucleotide base sequence that is identical or complimentary to a segment of 25 or 50 or 100 contiguous nucleotide bases of SEQ ID NO:23, 24 or 25, wherein the polynucleotide hybridizes to a polynucleotide that encodes a single chain variable region polypeptide that binds to a CD19 antigen.

In yet another embodiment, the present invention contemplates an isolated and purified single chain variable region polypeptide that binds to a CD19 antigen. Preferably, the polypeptide of this embodiment has a molecular weight of approximately 28 kDa. More preferably, the polypeptide binds to a CD19 antigen with a K_a of at least 1 x 109 M-1. More preferably still, the polypeptide comprises an amino acid residue sequence according to SEQ ID NO:20, 21 or 22.

In another aspect of this embodiment, the isolated and purified polypeptide of the claimed invention is further modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide. Also contemplated by the invention is a dimer of an isolated and purified single chain variable region polypeptide, wherein the dimer is prepared by linking a first polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide with a second polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide, the linking accomplished through a disulfide bond between a C-terminus cysteine residue on each polypeptide. In yet another embodiment, the present invention provides a process for preparing an isolated and purified single chain variable region polypeptide that binds to a CD19 antigen comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming *E. coli* cells with the expression vector; and (c) maintaining

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the transformed cells under biological conditions sufficient for expression of the polypeptide. Preferably, the process of this embodiment uses *E. coli* cells from a BL21(DE3) strain.

In another aspect, this embodiment of the invention provides an isolated and purified single chain variable region polypeptide that binds to a CD19 antigen, wherein the polypeptide is preparable by a process comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming *E. coli* cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide.

Preferably, this aspect of the invention provides an isolated and purified single chain variable region polypeptide that binds to a CD19 antigen, wherein the polypeptide is prepared by a process comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming *E. coli* cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide.

In an alternative aspect, the present invention contemplates a polypeptide prepared as described immediately above, wherein the polypeptide is further modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide. This aspect also provides a dimer of an isolated and purified single chain variable region polypeptide, wherein the dimer is prepared by linking a first polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide with a second polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide, the linking accomplished through a disulfide bond between a C-terminus cysteine residue on each polypeptide.

Also contemplated by the present invention is a polypeptide comprising an amino acid residue sequence of from five to sixty contiguous amino acid residues identical to any five to sixty contiguous amino acid residues of the polypeptide as defined by SEQ ID NO:20, 21 or

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22, wherein the polypeptide retains an ability to bind to a CD19 antigen with a K_a of at least 1 x 10^9 M⁻¹.

In an alternative embodiment, the claimed invention provides an immunoconjugate for the treatment of cancer comprising a single chain variable region polypeptide that binds to a CD19 antigen, wherein the polypeptide is linked to at least one cytotoxic agent. Preferably, the cancer susceptible to treatment with the immunoconjugate of the invention is a B-cell leukemia. More preferably, the immunoconjugate of this embodiment of the invention comprises an amino acid residue sequence according to SEQ ID NO:20, 21 or 22. Alternatively, the cytotoxic agent of the immunoconjugate of the invention is selected from the group consisting of single chain, double chain, and multiple chain toxins. In another aspect, the cytotoxic agent of the immunoconjugate of the invention is selected from the group consisting of beta-emitting metallic radionuclides, alpha emitters, and gamma emitters.

In yet another embodiment, the present invention provides an immunoconjugate for the treatment of cancer comprising a polypeptide that is a dimer of an isolated and purified single chain variable region polypeptide, wherein the dimer is prepared by linking a first polypeptide modified by the site specific insertion of a cysteine residue at the Cterminus of the polypeptide with a second polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide, the linking accomplished through a disulfide bond between a C-terminus cysteine residue on each polypeptide. and wherein the polypeptide is linked to at least one cytotoxic agent. Preferably, this immunoconjugate of the invention is efficacious for the treatment of Bcell leukemia. More preferably, the cytotoxic agent of this immunoconjugate is selected from the group consisting of single chain, double chain, and multiple chain toxins. Alternatively, the cytotoxic agent is a radionuclide selected from the group consisting of beta-emitting metallic radionuclides, alpha emitters, and gamma emitters. In another aspect, the immunoconjugate comprises both a toxin and a radionuclide.

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The present invention also contemplates a process for preparing an immunoconjugate comprising a single chain variable region polypeptide that binds to a CD19 antigen, wherein the process comprises the steps of (1) preparing the polypeptide according to a method comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming E. coli cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide; (2) providing a suitable toxin; and (3) conjugating the polypeptide to the toxin. As contemplated by the present invention, the process also further comprises the step of labelling the immunoconjugate with a radionuclide. Preferably, the toxin used in the process of the invention is selected from the group consisting of single chain, double chain, and multiple chain toxins. Likewise, the radionuclide used in the claimed process is selected from the group consisting of betaemitting metallic radionuclides, alpha emitters, and gamma emitters. In another aspect of this embodiment, the polypeptide of the immunoconjugate comprises an amino acid residue sequence according to SEQ ID NO:20, 21 or 22.

Alternatively, the present invention provides immunoconjugate for the treatment of cancer comprising a single chain variable region polypeptide that binds to a CD19 antigen, wherein the immunoconjugate is preparable by a process comprising the steps of (1) preparing the polypeptide according to a method comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming *E. coli* cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide; (2) providing a suitable toxin; and (3) conjugating the polypeptide to the toxin. Preferably, the immunoconjugate for the treatment of cancer comprising a single chain variable region polypeptide that binds to a CD19 antigen is prepared by a process comprising the steps of (1) preparing the polypeptide according to a method comprising the steps of (a) cloning a DNA sequence that encodes

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the polypeptide into an expression vector; (b) transforming *E. coli* cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide; (2) providing a suitable toxin; and (3) conjugating the polypeptide to the toxin.

Also provided is the immunoconjugate described immediately above, wherein the polypeptide of the immunoconjugate is linked to at least one cytotoxic agent. Preferably, the cytotoxic agent is selected from the group consisting of single chain, double chain, and multiple chain toxins or, alternatively, the cytotoxic agent is a radionuclide selected from the group consisting of beta-emitting metallic radionuclides, alpha emitters, and gamma emitters, or the immunoconjugate comprises one of each type of cytotoxic agent. Such an immunoconjugate is contemplated to be efficacious in the treatment of B-cell leukemia.

The present invention further contemplates an additional embodiment of a method for the treatment of cancer comprising the steps of (a) selecting a patient evidencing symptoms of a B-cell cancer, wherein the cancer is selected from the group consisting of leukemia and B-cell lymphoma; (b) administering to the patient, in a biocompatible dosage form, a therapeutically effective amount of an immunoconjugate, prepared according to a process comprising the steps of (1) preparing a polypeptide according to a method comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming E. coli cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide; (2) providing a suitable toxin; (3) conjugating the polypeptide to the toxin; and (4) labelling the immunoconjugate with a radionuclide. Preferably, the radionuclide with which the immunoconjugate is labelled is ¹³¹I.

In yet another embodiment, the present invention provides for a method for the treatment of cancer comprising the steps of (a) selecting a patient evidencing symptoms of a B-cell cancer, wherein the cancer is selected from the group consisting of leukemia and B-cell lymphoma; and

(b) administering to the patient, in a biocompatible dosage form, a therapeutically effective amount of an immunoconjugate comprising a polypeptide that is a dimer of an isolated and purified single chain variable region polypeptide, wherein the dimer is prepared by linking a first polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide with a second polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide, the linking accomplished through a disulfide bond between a C-terminus cysteine residue on each polypeptide. and wherein the polypeptide is linked to a toxin and labelled with a radionuclide.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1. Cloning strategy for development of anti CD19 scFv.

The variable domain of the heavy chain and the linker which encodes (G₄S)₃ were ligated into Bluescript K5 plasmid at *Xho1* and *Sac1* sites. Variable domains of the light chain were inserted into *Sst1* and *BgIII* sites following the linker. The pERT vector which was constructed by modifying pET3b was used as the expression vector for scFv. The nucleotides between *Ndc1* and *Xho1* sites of pERT encode four amino acids which are part of the FR1 of V_H but not included in the PCR products of V_H. The scFv encoding fragment was cloned into the pERT vector at *Xho1* and *Bgl II* sites. Positive clones were identified by restriction enzyme analysis and DNA sequencing.

Figure 2. Comparison of the DNA sequence of the different variable regions from the heavy and light chains (in two panels).

A: Heavy chain sequence. B: Light chain sequence. In the heavy chain CDR3, lower case letters are n nucleotide additions and they flank the germline encoded D_H gene sequences. Capitol letters indicate primers used in PCR.

Figure 3. Amino acid sequence alignment of the variable heavy and light chain regions from the three different hybridomas: B43, 25C1 and BLY3 (in two panels).

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Sequence differences are as indicated. The predicted protein sequences from the primers used for PCR are shown in bold type.

Figure 4. Expres

Expression and Purification of scFv.

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Lane 1, Molecular weight markers (97, 66, 45, 31, 21 KD); Lane 2, Uninduced cells; Lane 3, Induced cells; Lane 4, Sonicated supernatant; Lane 5, Detergent-solubilized supernatant; Lane 6, Pellet; Lane 7, Pellet purified by Q sepharose.

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Figure 5. Specific binding of FVS191 and FVS192 to CD19+ HLA Class I+ Cells in FACS.

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The X axis represents binding of FITC labelled class I antibody, Y axis represents binding of phycoerythrin labelled CD19 antibody. Panel A, negative control; panel B, positive control; panel C, specific blocking with FVS191; panel D, specific blocking with FVS192.

Figure 6. Scatchard analysis of binding of FVS191.

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Results are plotted with molecules/cell on horizontal axis and molecule L per cell mole on vertical axis. The derived K_a is 2 X 10^9 .

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DETAILED DESCRIPTION OF THE INVENTIO

There is a great need for the development of new therapeutic reagents for the treatment of a variety of diseases that are refractory to current therapies; one approach to developing these therapies has been through the use of monoclonal antibodies. The use of monoclonal antibodies in leukemia is particularly attractive because specific subsets of

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cells may be potentially specifically targeted. Several approaches have been tried using monoclonal antibodies for therapeutic use and often rely on the ability to chemically conjugate the antibodies to toxins (Ghetie *et al.*, 1988; Uckun *et al.*, 1986; Myers *et al.*, 1991; Jansen *et al.*, 1992). However, there are several disadvantages to use of intact antibodies particularly because of the large size of the molecules and the resultant relative inability to penetrate tissues (Pastan *et al.*, 1991; Yokota *et al.*, 1992).

Single chain fragments have been developed to overcome the problems associated with intact antibodies. scFvs contain only the variable regions from the heavy and light chains and have a molecular mass of approximately 28 kDa compared to that of the intact antibody of 150 kDa. However, many scFvs expressed in bacteria are insoluble, difficult to refold, and their ability to retain binding to the antigen of interest is highly variable. Because the effects of primary amino acid sequence on protein folding are not well understood, there is no known *a priori* method for determining the ability of a particular antibody to function when produced as an scFv. Accordingly, scFvs developed from three hybridomas that produce antibodies that bind to the CD19 antigen of B cells have been cloned and expressed.

20 Polynucleotides and Methods of the Invention.

In a first aspect, the present invention provides an isolated and purified polynucleotide encoding a single chain variable region polypeptide that binds to a CD19 antigen. Preferably, the isolated and purified polynucleotide of the invention encodes a polypeptide that has a molecular weight of approximately 28 kDa. More preferably, the polynucleotide the invention encodes a polypeptide that binds to a CD19 antigen with a K_a of at least 1 x 10⁹ M⁻¹. As used herein, the term "polynucleotide" means a sequence of nucleotides connected by phosphodiester linkages.

Also provided by the invention is a process for preparing an isolated and purified polynucleotide encoding a single chain variable region polypeptide that binds to a CD19 antigen comprising the steps of (a)

isolating RNA from hybridomas producing monoclonal antibodies to CD19 antigen; (b) transcribing isolated RNA to cDNA; (c) amplifying separate cDNA sequences encoding a heavy and a light variable region of the monoclonal antibody by a polymerase chain reaction; (d) cloning of separate amplification products encoding the heavy and the light variable regions, respectively, into vector constructs; (e) cloning of DNA encoding the heavy and the light variable regions, wherein the separate sequences are joined through a linker nucleotide sequence, into a vector construct; and (f) digesting the clones with appropriate restriction endonucleases. Preferably, the process of the invention utilizes a linker sequence that is the polynucleotide sequence described by SEQ ID NO:7.

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The present invention also contemplates an isolated and purified polynucleotide preparable by a process comprising the steps of (a) isolating RNA from hybridomas producing monoclonal antibodies to CD19 antigen; (b) transcribing isolated RNA to cDNA; (c) amplifying separate cDNA sequences encoding a heavy and a light variable region of the monoclonal antibody by a polymerase chain reaction; (d) cloning of separate amplification products encoding the heavy and the light variable regions, respectively, into vector constructs; (e) cloning of DNA encoding the heavy and the light variable regions, wherein the separate sequences are joined through a linker nucleotide sequence, into a vector construct; and (f) digesting the clones with appropriate restriction endonucleases.

Alternatively, the present invention provides an isolated and purified polynucleotide prepared by a process comprising the steps of (a) isolating RNA from hybridomas producing monoclonal antibodies to CD19 antigen; (b) transcribing isolated RNA to cDNA; (c) amplifying separate cDNA sequences encoding a heavy and a light variable region of the monoclonal antibody by a polymerase chain reaction; (d) cloning of separate amplification products encoding the heavy and the light variable regions, respectively, into vector constructs; (e) cloning of DNA encoding the heavy and the light variable regions, wherein the separate sequences are joined through a linker nucleotide sequence, into a vector construct;

and (f) digesting the clones with appropriate restriction endonucleases. Preferably, the process of the invention utilizes a linker sequence that is the polynucleotide sequence described by SEQ ID NO:7. More preferably, the isolated and purified polynucleotide of the claimed invention encodes a polypeptide comprising an amino acid residue sequence according to SEQ ID NO:20, 21 or 22. More preferably still, the isolated and purified polynucleotide of the invention comprises a nucleotide sequence according to SEQ ID NO:23, 24 or 25.

In an alternative embodiment, the present invention provides an isolated and purified polynucleotide comprising a nucleotide base sequence that is identical or complimentary to a segment of at least 10 contiguous nucleotide bases of SEQ ID NO:23, 24 or 25, wherein the polynucleotide hybridizes to a polynucleotide that encodes a single chain variable region polypeptide that binds to a CD19 antigen. Preferably, the polynucleotide of this embodiment of the invention encodes a polypeptide that has a molecular weight of approximately 28 kDa. More preferably still, the encoded polypeptide binds to a CD19 antigen with a K_a of at least 1 x 109 M-1.

In another aspect, this embodiment of the invention provides an isolated and purified polynucleotide comprising a nucleotide base sequence that is identical or complimentary to a segment of 25 or 50 or 100 contiguous nucleotide bases of SEQ ID NO:23, 24 or 25, wherein the polynucleotide hybridizes to a polynucleotide that encodes a single chain variable region polypeptide that binds to a CD19 antigen.

25 Polypeptides and Methods of the Invention.

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The scFv polypeptides developed from three hybridomas were expressed at high levels in bacteria. No instability of the protein, as determined by examination of Coomassie stained SDS-PAGE gels, was noted over the period of induction (3 hrs.) and all clones produced approximately the same quantities of protein. However, the ability of the scFv from each of these clones to bind to the target antigen varied greatly. Although the BLy3 and B43 hybridomas produced heavy chain and light

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chain variable proteins that were from the same family, only the protein produced from the B43 clone (FVS191) was able to show any ability to bind to the CD19 protein. This indicates the importance of the total sequence in the refolding of the native protein structure but indicates that development of scFv with proper folding and high binding affinity remains empiric. Like FVS191, the scFv clone from 25C1 (FVS192) also produced a protein capable of recognizing the antigen. However, the specific affinity of FVS192 for the CD19 antigen was low and could not be quantified in Scatchard analyses.

In yet another embodiment, the present invention contemplates an isolated and purified single chain variable region polypeptide that binds to a CD19 antigen. Preferably, the polypeptide of this embodiment has a molecular weight of approximately 28 kDa. More preferably, the polypeptide binds to a CD19 antigen with a K_a of at least 1 x 109 M-1. More preferably still, the polypeptide comprises an amino acid residue sequence according to SEQ ID NO:20, 21 or 22.

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In another aspect of this embodiment, the isolated and purified polypeptide of the claimed invention is further modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide. Also contemplated by the invention is a dimer of an isolated and purified single chain variable region polypeptide, wherein the dimer is prepared by linking a first polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide with a second polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide, the linking accomplished through a disulfide bond between a C-terminus cysteine residue on each polypeptide. As used herein the term "polypeptide" means a polymer of amino acids connected by amide linkages, wherein the number of amino acid residues can range from about 5 to about one million. Preferably, a polypeptide has from about 10 to about 1000 amino acid residues and, even more preferably from about 20 to about 500 amino residues. Thus, as used herein, a polypeptide includes what is often referred to in the art as an oligopeptide

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(5-10 amino acid residues), a polypeptide (11-100 amino acid residues) and a protein (>100 amino acid residues). A polypeptide encoded by an encoding region can undergo post-translational modification to form conjugates with carbohydrates, lipids, nucleic acids and the like to form glycopolypeptides (e.g., glycoproteins), lipopolypeptides (e.g. lipoproteins) and other like conjugates.

Polypeptides are disclosed herein as amino acid residue sequences. Those sequences are written left to right in the direction from the amino to the carboxy terminus. In accordance with standard nomenclature, amino acid residue sequences are denominated by either a single letter or a three letter code as indicated in Table 1 below.

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TABLE 1

Amino Acid Residue	3-Letter Code	1-Letter Code
Alanine	Ala	A
Arginine	Arg	R
Asparagine	Asn	N
Aspartic Acid	Asp	D
Cysteine	Cys	С .
Glutamine	Gln	Q
Glutamic Acid	Glu	E
Glycine	Gly	G
Histidine	His	Н
Isoleucine	Ile	I
Leucine	Leu	L
Lysine	Lys	K
Methionine	Met	M
Phenylalanine	Phe	F
Proline	Pro	P
Seriune	Ser	S
Threonine	Thr	T
Tryptophan	Trp	W
Tyrosine	Tyr	Y
Valine	Val	V
	Alanine Arginine Asparagine Aspartic Acid Cysteine Glutamine Glutamic Acid Glycine Histidine Isoleucine Leucine Lysine Methionine Phenylalanine Proline Seriune Threonine Tryptophan Tyrosine	Alanine Arg Asparagine Asn Aspartic Acid Asp Cysteine Cys Glutamine Gln Glutamic Acid Glu Glycine Gly Histidine His Isoleucine Ile Leucine Leu Lysine Lys Methionine Met Phenylalanine Phe Proline Seriune Ser Threonine Thr Tryptophan Trp Tyrosine Tyr

Modifications and changes may be made in the structure of a polypeptide of the present invention and still obtain a molecule having like characteristics. For example, certain amino acids can be substituted for other amino acids in a sequence without appreciable loss of activity. Because it is the interactive capacity and nature of a polypeptide that defines that polypeptide's biological functional activity, certain amino acid

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sequence substitutions can be made in a polypeptide sequence (or, of course, its underlying DNA coding sequence) and nevertheless obtain a polypeptide with like properties.

In making such changes, the hydropathic index of amino acids can be considered. The importance of the hydropathic amino acid index in conferring interactive biologic function on a polypeptide is generally understood in the art (Doolittle, et al. 1982). It is known that certain amino acids can be substituted for other amino acids having a similar hydropathic index or score and still result in a polypeptide with similar biological activity. Each amino acid has been assigned a hydropathic index on the basis of its hydrophobicity and charge characteristics.

Those indices are given in Table 2, below.

TABLE 2

15	Amino Acid	<u>Index</u>	Amino Acid	<u>Index</u>
	isoleucine	(+4.5)	tryptophan	(-0.9)
	valine	(+4.2)	tyrosine	(-1.3)
	leucine	(+3.8)	proline	(-1.6)
	phenylalanine	(+2.8)	histidine	(-3.2)
20	cysteine	(+2.5)	glutamate	(-3.5)
	methionine	(+1.9)	glutamine	(-3.5)
	alanine	(+1.8)	aspartate	(-3.5)
	glycine	(-0.4)	asparagine	(-3.5)
	threonine	(-0.7)	lysine	(-3.9)
25	serine	(-0.8)	arginine	(-4.5)

It is believed that the relative hydropathic character of the amino acid determines the secondary structure of the resultant polypeptide, which in turn defines the interaction of the polypeptide with other molecules, for example, enzymes, substrates, receptors, antibodies, antigens, and the like. It is known in the art that an amino acid may be substituted by another amino acid having a similar hydropathic index and still obtain a biologically functionally equivalent polypeptide. In such

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changes, the substitution of amino acids whose hydropathic indices are within \pm 2 is preferred, those which are within \pm 1 are particularly preferred, and those within \pm 0.5 are even more particularly preferred.

Substitution of like amino acids can also be made on the basis of hydrophilicity, particularly where the biologically functionally equivalent peptide or polypeptide thereby created is intended for use in immunological embodiments. U.S. Patent 4,554,101, incorporated herein by reference, states that the greatest local average hydrophilicity of a polypeptide, as governed by the hydrophilicity of its adjacent amino acids, correlate with its immunogenicity and antigenicity, *i.e.* with a biological property of the polypeptide.

As detailed in U.S. Patent 4,554,101, the following hydrophilicity values have been assigned to amino acid residues: arginine (+3.0); lysine (+3.0); aspartate (+3.0 \pm 1); glutamate (+3.0 \pm 1); serine (+0.3); asparagine (+0.2); glutamine (+0.2); glycine (0); proline (-0.5 \pm 1); threonine (-0.4); alanine (-0.5); histidine (-0.5); cysteine (-1.0); methionine (-1.3); valine (-1.5); leucine (-1.8); isoleucine (-1.8); tyrosine (-2.3); phenylalanine (-2.5); tryptophan (-3.4). It is understood that an amino acid can be substituted for another having a similar hydrophilicity value and still obtain a biologically equivalent, and in particular, an immunologically equivalent, polypeptide. In such changes, the substitution of amino acids whose hydrophilicity values are within \pm 2 is preferred, those which are within \pm 1 are particularly preferred, and those within \pm 0.5 are even more particularly preferred.

As outlined above, amino acid substitutions are generally therefore based on the relative similarity of the amino acid side-chain substituents, for example, their hydrophobicity, hydrophilicity, charge, size, and the like. Exemplary substitutions which take various of the foregoing characteristics into consideration are well known to those of skill in the art and include: arginine and lysine; glutamate and aspartate; serine and threonine; glutamine and asparagine; and valine, leucine and isoleucine. The present invention thus contemplates functional equivalents of the

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claimed polypeptides.

In yet another embodiment, the present invention provides a process for preparing an isolated and purified single chain variable region polypeptide that binds to a CD19 antigen comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming *E. coli* cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide. Preferably, the process of this embodiment uses *E. coli* cells from a BL21(DE3) strain.

In another aspect, this embodiment of the invention provides an isolated and purified single chain variable region polypeptide that binds to a CD19 antigen, wherein the polypeptide is preparable by a process comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming *E. coli* cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide.

Preferably, this aspect of the invention provides an isolated and purified single chain variable region polypeptide that binds to a CD19 antigen, wherein the polypeptide is prepared by a process comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming *E. coli* cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide.

In an alternative aspect, the present invention contemplates a polypeptide prepared as described immediately above, wherein the polypeptide is further modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide. This aspect also provides a dimer of an isolated and purified single chain variable region polypeptide, wherein the dimer is prepared by linking a first polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide with a second polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide, the

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linking accomplished through a disulfide bond between a C-terminus cysteine residue on each polypeptide.

Also contemplated by the present invention is a polypeptide comprising an amino acid residue sequence of from five to sixty contiguous amino acid residues identical to any five to sixty contiguous amino acid residues of the polypeptide as defined by SEQ ID NO:20, 21 or 22, wherein the polypeptide retains an ability to bind to a CD19 antigen with a K_a of at least 1 x 109 M-1.

Immunoconjugates and Methods of the Invention.

In an alternative embodiment, the claimed invention provides an immunoconjugate for the treatment of cancer comprising a single chain variable region polypeptide that binds to a CD19 antigen, wherein the polypeptide is linked to at least one cytotoxic agent. Preferably, the cancer susceptible to treatment with the immunoconjugate of the invention is a B-cell leukemia. More preferably, the immunoconjugate of this embodiment of the invention comprises an amino acid residue sequence according to SEQ ID NO:20, 21 or 22. Alternatively, the cytotoxic agent of the immunoconjugate of the invention is selected from the group consisting of single chain, double chain, and multiple chain toxins. In another aspect, the cytotoxic agent of the immunoconjugate of the invention is selected from the group consisting of beta-emitting metallic radionuclides, alpha emitters, and gamma emitters.

Toxins

A structural similarity in plant and bacterial toxins inhibits protein synthesis: they are usually heterodimers made of a polypeptide chain (B chain) that binds the toxin to target cells and a second chain (A chain) that displays enzymatic activity (Olsnes *et al.*, 1982). The two chains are linked by a disulfide bond. Diphtheria toxin is a slight exception in that a single proteolytic cleavage is required to generate an A and a B chain (Collier *et al.*, 1971) that are also disulfide bonded. In addition, it is provocative that the subunits of all the plant toxins have approximately the same apparent molecular weight (Olsnes *et al.*, 1982; Olsnes *et al.*, 1974), about 30,000, that

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the A chains attack the 60S ribosomal subunit (Olsnes et al., 1982; Olsnes et al., 1974; Olsnes et al., 1984) and the B chains bind to galactose (Olsnes et al., 1982; Olsnes et al., 1974; Olsnes et al., 1984). Moreover, the A and B chains of abrin and ricin, two toxins derived from phylogenetically distant plants, can be interchanged to produce hybrid molecules of relatively high toxicity (Olsnes et al., 1982; Olsnes et al., 1984). These observations suggest significant conservation in function and structure. Whether the structural conservation is at the three-dimensional level only or reflects primary amino acid sequence homologies remains to be determined. There is also a variety of plant toxins composed of A chains only, e.g., gelonin (Stirpe et al., 1980) and pokeweed antiviral protein (PAP) (Olsnes et al., 1982; Barbieri et al., 1982). These A chains function in the same way as the A chains of intact toxins.

In yet another embodiment, the present invention provides an immunoconjugate for the treatment of cancer comprising a polypeptide that is a dimer of an isolated and purified single chain variable region polypeptide, wherein the dimer is prepared by linking a first polypeptide modified by the site specific insertion of a cysteine residue at the Cterminus of the polypeptide with a second polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide, the linking accomplished through a disulfide bond between a C-terminus cysteine residue on each polypeptide. and wherein the polypeptide is linked to at least one cytotoxic agent. Preferably, this immunoconjugate of the invention is efficacious for the treatment of Bcell leukemia. More preferably, the cytotoxic agent of this immunoconjugate is selected from the group consisting of single chain, double chain, and multiple chain toxins. Alternatively, the cytotoxic agent is a radionuclide selected from the group consisting of beta-emitting metallic radionuclides, alpha emitters, and gamma emitters. In another aspect, the immunoconjugate comprises both a toxin and a radionuclide.

The present invention also contemplates a process for preparing an immunoconjugate comprising a single chain variable region polypeptide

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that binds to a CD19 antigen, wherein the process comprises the steps of (1) preparing the polypeptide according to a method comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming E. coli cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide; (2) providing a suitable toxin; and (3) conjugating the polypeptide to the toxin. As contemplated by the present invention, the process also further comprises the step of labelling the immunoconjugate with a radionuclide. Preferably, the toxin used in the process of the invention is selected from the group consisting of single chain, double chain, and multiple chain toxins. Likewise, the radionuclide used in the claimed process is selected from the group consisting of betaemitting metallic radionuclides, alpha emitters, and gamma emitters. In another aspect of this embodiment, the polypeptide of the immunoconjugate comprises an amino acid residue sequence according to SEQ ID NO:20, 21 or 22.

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The toxins which are usable in the practice of the claimed invention encompass all toxins used in the production of immunotoxins. Generally, the toxins include heterodimers made of a polypeptide chain (B chain) that binds the toxin to target cells via a sugar on the surface and a second chain (A chain) that displays enzymatic activity. The two chains are typically linked by a disulfide bond. Examples of two chain toxins are ricin, abrin, modeccin, diphtheria toxin and viscumin. However, single chain toxins, *i.e.* toxins composed of A chains only, *e.g.*, gelonin, pseudomonas aeruginosa Exotoxin A, and amanitin may also be utilized. Other single chain toxins are hemitoxins which are also usable in this invention. They include pokeweed antiviral protein (PAP), saporin and memordin. Other useful single chain toxins include the A-chain fragments of the two chain toxins. A chain toxins with multiple B chains such as Shigella toxin are also usable in the invention.

As used herein, 2-chain toxins refers to toxins formed from two chains, and single chain toxins refers to both toxin obtained by cleaving 2-

chain toxins as well as toxins having only one chain.

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A preferred toxin is ricin, a toxin lectin extracted from the seeds of Ricinus communis, which contains an enzymatic and protein synthesis inhibiting A chain and a B chain which contains galactose binding site(s). Ricin is extremely toxic and it has been calculated that a single molecule of ricin in the cytosol will kill a cell. Ricin may be obtained and purified by the procedures described in U.S. Pat. No. 4,340,535, the disclosure of which is incorporated herein by reference.

Alternatively, the present invention provides an immunoconjugate for the treatment of cancer comprising a single chain variable region polypeptide that binds to a CD19 antigen, wherein the immunoconjugate is preparable by a process comprising the steps of (1) preparing the polypeptide according to a method comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming E. coli cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide; (2) providing a suitable toxin; and (3) conjugating the polypeptide to the toxin. Preferably, the immunoconjugate for the treatment of cancer comprising a single chain variable region polypeptide that binds to a CD19 antigen is prepared by a process comprising the steps of (1) preparing the polypeptide according to a method comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming E. coli cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide; (2) providing a suitable toxin; and (3) conjugating the polypeptide to the toxin. One general method of preparing immunotoxins is to use a thiolcontaining heterobifunctional crosslinker, e.g., SPDP, which attacks primary amino groups on the antibody and by disulfide exchange can attach either the SH-containing A chain or the SPDP-derivatized holotoxin to the antibody (Cumber et al., 1984; Carlsson et al., 1978). If the disulfide exchange is carried out at neutral pH a relatively stable disulfide bond is

formed and the conjugate remains intact when incubated with fresh mouse serum *in vitro*.

The nature of the linkage between the A chain and the antibody or fragment is of critical important in determining toxicity. If the bond cannot be broken readily in an endosome/phagolysosome (Jansen *et al.*, 1982; Ramakrishnan *et al.*, 1984), *e.g.*, a stable thioether bond, then toxicity is virtually abolished (Jansen *et al.*, 1982). In contrast, if the bond is highly unstable, then the conjugate may dissociate either before it reaches the target cell or, perhaps, prematurely within the target cell. In the latter case, the A chain may be degraded before translocation can occur.

Also provided is the immunoconjugate described above, wherein the polypeptide of the immunoconjugate is linked to at least one cytotoxic agent. Preferably, the cytotoxic agent is selected from the group consisting of single chain, double chain, and multiple chain toxins or, alternatively, the cytotoxic agent is a radionuclide selected from the group consisting of beta-emitting metallic radionuclides, alpha emitters, and gamma emitters, or the immunoconjugate comprises one of each type of cytotoxic agent. Such an immunoconjugate is contemplated to be efficacious in the treatment of B-cell leukemia.

Among the radionuclides used, gamma-emitters, positron-emitters, and X-ray emitters are suitable for localization and/or therapy, while beta emitters and alpha emitters may also be used for therapy. Suitable radionuclides for forming the immunoconjugate of the invention include 123I, 125I, 130I, 131I, 133I, 135I, 47Sc, 72As, 72Se, 90Y, 88Y, 97Ru, 100Pd, 101mRh, 119Sb, 128Ba, 197Hg, 211At, 212Bi, 212Pb, 109Pd, 111In, 67Ga, 68Ga, 67Cu, 75Br, 77Br, 99mTc, 11C, 13N, 15O and 18F.

Methods for the Treatment of Cancer.

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The present invention further contemplates an additional embodiment of a method for the treatment of cancer comprising the steps of (a) selecting a patient evidencing symptoms of a B-cell cancer, wherein the cancer is selected from the group consisting of leukemia and B-cell lymphoma; (b) administering to the patient, in a biocompatible dosage

form, a therapeutically effective amount of an immunoconjugate, prepared according to a process comprising the steps of (1) preparing a polypeptide according to a method comprising the steps of (a) cloning a DNA sequence that encodes the polypeptide into an expression vector; (b) transforming *E. coli* cells with the expression vector; and (c) maintaining the transformed cells under biological conditions sufficient for expression of the polypeptide; (2) providing a suitable toxin; (3) conjugating the polypeptide to the toxin; and (4) labelling the immunoconjugate with a radionuclide. Preferably, the radionuclide with which the immunoconjugate is labelled is ¹³¹I.

In yet another embodiment, the present invention provides for a method for the treatment of cancer comprising the steps of (a) selecting a patient evidencing symptoms of a B-cell cancer, wherein the cancer is selected from the group consisting of leukemia and B-cell lymphoma; and (b) administering to the patient, in a biocompatible dosage form, a therapeutically effective amount of an immunoconjugate comprising a polypeptide that is a dimer of an isolated and purified single chain variable region polypeptide, wherein the dimer is prepared by linking a first polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide with a second polypeptide modified by the site specific insertion of a cysteine residue at the C-terminus of the polypeptide, the linking accomplished through a disulfide bond between a C-terminus cysteine residue on each polypeptide, and wherein the polypeptide is linked to a toxin and labelled with a radionuclide.

25 EXAMPLES

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Example 1: Cloning and Expression of the scFv

A. Cloning of the variable regions (V_H and V_L)

<u>Cells</u>: The three anti-CD19 hybridomas used in these studies have been previously described: B43, produced by F. Uckun (Uckun *et al.*, 1986), SJ25C1, produced by S. Pieper, and BLY3, produced by S. Poppema (Knapp *et al.*, 1989b). All were maintained in RPMI 1640 supplemented with 10% fetal calf serum.

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RNA was isolated by the method of Chomczynski and Sacchi (Chomczynski, 1987) and either used directly for RT-PCR or further purified by oligo dT column chromatograph. By way of example, and without limitation, the following protocol describes isolation of RNA from 100 mg of rat mammary tissue according to the method referenced above.

Immediately after removal from the animal, the tissue was minced on ice and homogenized (at room temperature) with 1 ml of solution D in a glass-Teflon homogenizer and subsequently transferred to a 4-ml polypropylene tube. Sequentially, 0.1 ml of 2M sodium acetate, pH 4, 1 ml of phenol (water saturated), and 0.2 ml of chloroform-isoamyl alcohol mixture (49:1) were added to the homogenate, with thorough mixing by inversion after the addition of each reagent. The final suspension was shaken vigorously for 10 s and cooled on ice for 15 min. Samples were centrifuged at 10,000g for 20 min. at 4°C. After centrifugation, RNA was present in the aqueous phase whereas DNA and proteins were present in the interphase and phenol phase. The aqueous phase was transferred to a fresh tube, mixed with 1 ml of isopropanol, and then placed at -20°C for at least 1 h to precipitate RNA. Sedimentation at 10,000g for 20 min. was again performed and the resulting RNA pellet was dissolved in 0.3 ml of solution D, transferred into a 1.5-ml Eppendorf tube, and precipitated with 1 vol of isopropanol at -20°C for 1 h. After centrifugation in an Eppendorf centrifuge for 10 min. at 4°C the RNA pellet was resuspended in 75% ethanol, sedimented, vacuum dried (15 min.), and dissolved in 50 µl 0.5% SDS at 65°C for 10 min. At this point the RNA preparation could be used for poly(A)+ selection by oligo (dT) chromatography, Northern blot analysis, and dot blot hybridization. Isopropanol precipitation can be replaced by precipitation with a double volume of ethanol.

Reverse transcription of the isolated RNA was performed according to the recommendations of the manufacturer (Life Technologies) using random hexamers and was performed in a 50 microliter reaction volume with 1-2 micrograms of polyadenylated RNA or 5-10 micrograms of total

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RNA. Approximately 10 microliters of the reverse transcribed material was used for the polymerase chain reaction using one pair of the several different primers listed in Table 1. The primers Z221 and Z222 anneal to the constant regions of heavy and light chains, respectively, and were only used for isolating clones for verification of sequence but not for the production of variable regions that were subsequently used in the creation of the scFv. The cycle parameters were 1 cycle of 94°C for 5' before the addition of the TaqI polymerase then 30 cycles of 94° C 1' 30", 54°C 1' 30", 72°C 1', followed by 1 cycle of 94°C 1'30", 54°C 2'30", 72°C 10'. The PCR products were cloned either after treatment with Klenow into Smal digested pBluescript or directly using the pCRI vector (Invitrogen) which has compatible T overhangs. Clones were identified based on the size of inserts (approximately 350bp for the V_L gene and 450bp for the V_H gene) and were confirmed by sequencing using standard dideoxynucleotide chain termination techniques (Sequenase, US Biochemicals). At least three different clones from three different PCR reactions were sequenced for each variable region to confirm the absence of any mutations induced by *Taq* polymerase before clones were used for the creation of scFV.

The DNA and the predicted amino acid sequences of the clones of the variable regions from the three hybridomas are shown in Fig. 2 and Fig. 3. As discussed in Materials and Methods, at least three clones from three independent PCR reactions were sequenced to ensure that no Taq-introduced mutations were present within the clones that were used for the scFv development. All heavy chain variable regions from the three hybridomas were from the J558 family which includes approximately 50% of all mouse heavy chain variable region genes (Brodeur *et al.*, 1984). Although clone 25CI uses J_H2, clones of B43 and BLy3 use J_H4. As expected, the B43 and Bly3 clones differed most within the CDR3 region due to N region differences.

Sequencing of the light chain variable regions showed that V_K21 was used in both B43 and Bly3 but V_K19 was used in 25C1. The J_K regions used were J_K1 for B43 and Bly3 and J_K2 for 25C1 (Sakano *et al.*, 1979). As

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anticipated, the greatest region of variability was present in the CDR3 region due to differential splicing and N region additions. After clones without any apparent PCR-introduced mutations had been identified by sequence analysis, scFvs were constructed.

B. Cloning of the scFv using V_H and V_L .

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The linker used in these studies was (Gly₄Ser)₃ as previously described (Huston *et al.*, 1988). The scFvs were created by ligation of the linker region oligonucleotides (Table 3) using the strategy outlined in FIG. 1. Heavy chain variable region was mixed simultaneously with linker and Bluescript to obtain the V_H-linker construct shown in FIG. 1. Clones that contained the heavy chain variable region were digested with *XhoI* and *BstEII*. Success of the procedure was confirmed by sequencing. Clones that contained the heavy chain variable region and the linker were then digested with *SstI* and *BglII* and ligated to gel purified light chain variable region that was digested with the same enzymes. Clones were identified by the appearance of appropriately sized restriction endonuclease fragments and finally by nucleotide sequence analysis. scFvs were then digested with *XhoI* and *BglII* and gel purified before ligation into the pERT expression vector.

TABLE 3. OLIGONUCLEOTIDES USED FOR scF_V CONSTRUCT Primer Name Oligonucleotide Sequence 5' VH: Z462 AGGTCCAGCTGCTCGAGTCTGG Xho1 TGAGGAGACGGTGACCGTGTCCCTTGGCCCCAG 3' VH: B1867 Ι **BstEII** 3' VH: Z221 AGGCTTACTAGTACAATCCCTGGGCACAAT 5' VK: Z407 CGCGGATCCAGTTCCGAGCTCGTGCTCACCCAGTCTCCA Ι Sst1 3' VK: B1865 GAAGATCTACGTTTTATTTCCAGCTTGGTCCC Ι Bgl1 3' VK: Z222 GCGCCGTCTAGAATTAACACTCATTCCTGTTGAA Linker for V_H and GGAG GCGGTGG CTCGGGC GGTGGCG GCTCGGG TGGCGGC GGAT CC V_L

The primers Z221, 222, 407 and 462 are based on sequences from Huse et al. (1989). The primers B1867 adn 1865 are based on primer sequences from Orlandi *et al.* (1989). The * denotes primers that were used for the generation of clones used only for sequencing. The oligonucleotides used for the liner are based on the three developed by Huston *et al.*

C. Expression of scFvs.

The vector used to express the scFv in these studies was developed using the pET3b plasmid established by Studier *et al.* (Studier *et al.*, 1990). This plasmid vector was developed for cloning and expressing target

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DNAs under control of a T7 promoter and designated pET vectors (plasmid for expression by T7 RNA polymerase) (Rosenberg *et al.*, 1987). These vectors contain a T7 promoter inserted into the BamHI site of the multi-copy plasmid pBR322 in the orientation that transcription is directed counterclockwise, opposite to that from the TET promoter. In the absence of T7 RNA polymerase, transcription of target DNAs by *E. coli* RNA polymerase is low enough that very toxic genes can be cloned in these vectors. However, some expression can be detected, so it is possible that an occasional gene may be too toxic to be cloned in them.

Most of the pET vectors described confer resistance to ampicillin. In such vectors, the *bla* gene is oriented so that it will be expressed from the T7 promoter along with the target gene. However, in the pET-9 series of vectors, the *bla* gene has been replaced by *kan* gene in the opposite orientation. In these vectors, the only coding sequence transcribed from the T7 promoter is that of the target gene.

The T7 promoter in the pET vectors is derived from the $\emptyset10$ promoter, one of six strong promoters in T7 DNA that have the identical nucleotide sequence from positions -17 to +6, where +1 is the position of the first nucleotide of the RNA transcribed from the promoter. The $\emptyset10$ promoter fragments carried by the vectors all begin at bp -23 and continue to bp +2, +3, +26, and +96 or beyond. Some of the vectors also contain a transcription termination signal or an RNase III cleavage site downstream of the cloning site for the target DNA.

pET3b was modified to allow for the cloning and expression of the constructs of the present invention by ligating an oligonucleotide that coded for the first four amino acids (LESG) that are commonly found at the amino terminus of the heavy chain variable region to the vector that was digested with NdeI and EcoRI. This oligonucleotide also contained sequences for recognition sites for XhoI, BgIII, BamHI, and EcoRI allowing for the cloning of the scFv into the vector at the XhoI and BgIII sites with the possibility of cloning other potentially therapeutic genes in the future (Table 3).

Expression of protein was accomplished by introducing the scFv clones into either BL21(DE3) or BL21(DE3) pLysS *E.coli* cells. No difference in the amount of recombinant protein expressed by these host strains was observed. Induction of protein synthesis was performed with 1.0 mM IPTG for three hours prior to harvesting of cells. Pellets were boiled in SDS-Page loading buffer and subjected to electrophoresis in denaturing polyacrylamide gels (Laemmli, 1970). Bacteria could be successfully induced at an O.D. 600 nm. of 0.6-1.0 if grown in a standard Erlenmeyer flask or at an O.D. 600 nm. of 2.5-3.0 if grown in a Fernbach (baffled) flask. Although the amount of protein per cell did not appear to change between cells grown in either flask as determined by SDS-PAGE and Coomassie staining (data not shown) the total amount of protein was greater from cells grown in the baffled flasks due to their greater mass.

Constructs were used to direct the synthesis of protein in *E. coli* as described above. After induction the protein was subjected to SDS-PAGE and detected by staining with Coomassie brilliant blue (Fig. 4). The results show that a protein of the expected molecular weight (27.5kDa) was specifically induced by the addition of IPTG to the culture medium. This protein was not present in either control cells or cells that were not treated with IPTG.

Example 2: Isolation of Protein.

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The isolation procedure for the scFvs followed that of previously published methods (Langley *et al.*, 1987). As described below, all of the protein produced was found within the insoluble cytoplasmic fraction presumably in inclusion bodies. Briefly, cells were harvested by centrifugation and washed in water before being resuspended in up to 1/5 of the original culture volume of 10 mM Tris-HCl pH 7.4, 50 mM NaCl. If the original culture volume was large (greater than 100mls) this solution was frozen at -20° C to ensure full lysis of the cells. The mixture was sonicated and centrifuged at 30,000x g for 30 minutes. The supernatant was discarded and the pellet was resuspended by sonication in 1/20 of the original culture volume in 10 mM Tris-HCl pH 8.0 and 5 mM EDTA.

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After resuspension was complete the mixture was digested with 0.2% lysozyme (Sigma) for a minimum of 1hour. Finally, 1/3 volume of 10% sodium deoxycholate was added and the mixture was incubated at room temperature for 1 hour before centrifugation at 30,000 x g for 30 minutes. The pellet was washed three times in water by resuspending the pellet by sonication and centrifugation as described above. Pellets were either stored at -20° C or dissolved in 0.1M Tris-HCl ph 8.0, 6M guanidine HCl, 0.3M DTE, 2mM EDTA at room temperature for a minimum of 2hours. Refolding of the denatured scFv was performed according to the method of Buchner *et. al.* (15).

The protein concentration was measured using the Bradford assay and the solution was then rapidly diluted at least 100 fold to a concentration of 30 ug/ml protein in 0.1M Tris-HCl pH8.0, 0.5M Larginine, 8mM GSSG and 2mM EDTA. After a minimum of 12 hours, at 10° C the refolded protein was concentrated using an Amicon spiral concentrator and spin concentrator before being chromatographed on Q Sepharose and finally Superose 75. As judged by the presence of a single peak on Superose chromatography and Coomassie stain of SDS-Page gels, protein was pure. If the concentration was too low for use in experiments the protein was concentrated by Amicon spin concentrators. Concentration of the protein was determined using the Bradford assay (BioRad) with bovine serum albumin as a standard.

To determine if the protein was present in an insoluble or soluble fraction, cells were disrupted by sonication and the supernatant and insoluble material separated by centrifugation. Analysis of the two fractions indicated that the bulk, if not all of the protein was present in the insoluble pellet. Due to the insolubility of the protein and its probable location within inclusion bodies, we performed isolations based on previously published methods for the purification of proteins from these vesicles. Refolded and purified protein was then used for FACS and Scatchard analysis.

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Example 3: Analysis of the scFvs

I. <u>FACS Analysis</u>.

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FACS analysis was performed on either RS4:11 (Stong *et al.*, 1985) or B1 cell line (Cohen *et al.*, 1991), both of which express CD19 and HLA Class I and carry the 4:11 translocation.

The RS4:11 cell line was established from bone marrow of a patient with t(4:11)-associated acute leukemia. Morphological, immunologic, and cytochemical characteristics of RS4:11 cells were found to be consistent with ALL. The cells are strongly positive for TdT. An in-depth analysis of RS4:11 revealed characteristics of both lymphoid and myeloid lineages.

The cells are rearranged for immunoglobulin heavy and k-chain genes, providing strong evidence for a commitment to B cell lineage. Although occasional heavy chain gene rearrangements have been noted in T cells and myeloid cells, light chain gene rearrangements have been restricted to the B cell lineage (Arnold et al., 1983; Korsmeyer et al., 1983; Ford et al., 1983). The expression of B4 is additional support for B lineage classification, since within the hematopoietic system, this antigen is expressed very early in normal B cell ontogeny and is restricted to B lineage cells (Nadler et al., 1983). Reactivity with BA-1, BA-2, and PI153/3 is consistent with B lineage classification because these MoAbs react with normal pre-B and B cells as well as with the vast majority of non-T ALL, although their binding cannot be considered to be definitive for lymphoid leukemias (LeBien et al., 1983).

In addition to these lymphoid characteristics, RS4:11 cells bind 1G10, a mAb that reacts with granulocytic cells, some monocytes (Bernstein *et al.*, 1988), and CFU-GM precursor cells (Andrews *et al.*, 1983). Some RS4:11 cells weakly express the gp170,95/TA-1 antigen found on monocytic precursors (Andrews *et al.*, 1983) and peripheral blood monocytes (LeBein *et al.*, 1980). The ultrastructural detection of basophil/mast cell granules and peroxidase activity in a minor population of RS4:11 cells is supportive evidence of myeloid commitment. Similar basophil/mast cell granules have been detected in some cases of lymphoid blast crisis of chronic

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myelogenous leukemia and in Philadelphia-positive ALL (Parkin *et al.*, 1982). The disappearance of this more differentiated subpopulation of RS4:11 suggests that these cells were at proliferative disadvantage or that the in vitro conditions could not support their phenotypic expression.

The monocyte-like phenotype of RS4:11 induced after TPA treatment is persuasive evidence for the myelomonocytic nature of RS4:11. Several laboratories have reported that TPA can induce human myeloid and lymphoid leukemic cells to more differentiated phenotypes that are primarily dictated by the differentiative potential of the target cells (Koeffler, 1983; LeBien et al., 1982; Nadler et al., 1982; Nagasawa et al., 1980). In response to TPA (0.5 to 10.0 ng/mL), RS4:11 cells became reactive with TA-1, OKM1, and MCS2, became phagocytic, and showed greatly enhanced NSE activity in a pattern characteristic of monocytic cells. subpopulation of treated cells became adherent, but this response resembled the weak adherence of certain TPA-treated lymphoid lines (Castagna et al., 1979) more closely than the strong adherence displayed by treated myeloid lines, such as HL-60 and KG-1 (Koeffler, 1983; Goodwin et Ultrastructurally, treated cells exhibited a monocytoid al., 1984). morphology.

The cell line B1 was established from bone marrow obtained from a 14-year-old child in first relapse. The patient's bone marrow sample at diagnosis and relapse contained over 95% malignant cells characterized by the t(4:11) (q21;q23) chromosomal translocation and biphenotypic expression of lymphoid and myeloid cell markers (often associated with this translocation).

The cell line was established by incubating leukemic cells ($10^6/\text{mL}$) in ∞ -MEM containing 10% heat-inactivated fetal calf serum (FCS). After 8 weeks, the cells were cloned in semisolid methylcellulose and single colonies were isolated and expanded in liquid culture medium. The cell line established this way resembled the donor's leukemic cells. The karyotype of the line showed t (4:11) (q21; a23) in all metaphases. In addition, other chromosomal abnormalities, including trisomy 6,

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der(1)t(1;8) (p36; q13), der(10)t(1;10)(q11; p15), were consistently observed in all metaphases. Cytochemical analysis showed a profile of periodic acid Schiff (PAS)-positive, acid phosphatase-positive, nonspecific esterase-positive, and Sudan black-negative staining. The leukemic cells lacked T-and B-cell markers (E-, sIg-, cIg-) and were CD10- and CD20-, but had undergone IgH(μ) gene rearrangement. Flow cytometric analysis showed that B1 cells expressed early pre-B-cell markers such as CD19+ and HLA-DR+. HLA-DR is coexpressed with My-9 (CD33), a marker of myeloid lineage on 20% of the cells. Other myeloid differentiation markers, such as My-7, Mo-1, and Mo-2, were undetectable on the surface of B1 cells.

These differentiation markers expressed on the B1 cell line are consistent with the early B and myeloid biphenotypic nature of the original bone marrow cells from this patient at relapse, and with previous reports of the association of the 4:11 translocation with biphenotypic leukemia.

All reactions were carried out at 4° C. Cells were counted and approximately 105 cells were aliquoted into polystyrene tubes. The cells were then incubated with FACS buffer (PBS containing 1% calf serum) for 20 minutes to block non-specific adherence of the antibodies. Cells were stained with the antibodies or scFv in a total volume of 200 µl for 20 minutes before being centrifuged and the supernatant discarded. Cells were then washed twice with FACS buffer before addition of 200 µl of biotinylated 25C1 antibody and streptavidin conjugated to phycoerythrin. The antibody was removed and the cells were washed again before addition of the anti HLA-class I antibody conjugated to FITC. After a final series of washes the cells were resuspended in PBS containing 0.4% paraformaldehyde. Fluorescence staining was measured by flow cytometry.

FACS analyses were used to evaluate the scFvs. The scFvs from B43 and 25C1 hybridomas (which are refered to as FVS191 (Fragment, Variable, Single chain, anti CD19, number 1) and FVS192, respectively, were able to inhibit the binding of FITC labeled 25C1 but not an anti HLA class I

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monclonal antibody to cells that were CD19+, HLA class 1+ (Fig. 5). The scFv derived from BLy3 (FV S193) did not block the binding of the competing antibody. Also, binding to target cells could not be detected by biotinylating the scFv developed from this hybridoma and using this material with streptavidin labeled phycoerythrin (data not shown). The failure of BLy3 scFv to bind in these two assays suggests that the protein was not properly folded.

II. <u>Scatchard analysis</u>.

Iodine labelling of the proteins was accomplished using Iodobeads (Pierce) and the specific activity was determined. Beads were washed with iodination buffer, dried, and added to solution of carrier free Na125I (1 mCi/100 μg of protein) and allowed to react for five minutes. The reaction was stopped and the beads were washed. Gel filtration (Pharmacia PD5) was used to remove excess Na125I. TCA precipitation was carried out followed by determination of specific activity using standard calculations. Immunoreactive fractions were subsequently determined (with reagents generally in the range of 0.05). Scatchard analysis was determined using FACS buffer and labelled protein diluted serially in unlabelled protein to give a final concentration of 200 $\mu g/mL$. Iodinated protein was purified by Dowex or size exclusion column chromatography and the specific activity was calculated.

Due to the ability of FVS191 and FVS192 to specifically bind to cells that express the CD19 antigen we evaluated their affinity. Proteins were iodinated and used for Scatchard analysis as described in Materials and Methods. The results (Fig.6) demonstrated that the FVS191 had an K_a of 2x10⁹ M⁻¹. Although FVS192 was able to successfully compete with 25C1 binding to the CD19 antigen it did not demonstrate sufficient avidity of binding to be evaluated in Scatchard analysis and its Ka therefore could not be determined.

30 Example 4: Formation of dimers of Anti-CD19 Single-Chain Fv.

Single-chain Fv antibody fragments have the advantage of improved tumor penetration over intact antibody. Dimers of scFv may

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possess higher binding constants and have potential as diagnostic or therapeutic agents.

To facilitate dimer formation, an additional cysteine residue was site-specifically inserted at the C-terminal of the scFv constructs of the present invention to form the scFv-cys. The scFv-cys proteins were isolated from bacterial inclusion bodies, reduced with guanidine, and refolded in redox buffer containing DTE and GSSG. Q-Sepharose-purified scFv-cys proteins were treated with 2 mM DTT. The DTT was removed using a Pharmacia PD10 column. Disulfide bonds between C-terminal cysteins were formed by air oxidation. Dimer formation of both B43 scFv-cys and 25C1 scFv-cys was confirmed by reducing and non-reducing SDS-PAGE. The scFv without C-terminal cysteine did not form covalently-linked dimers under these conditions, indicating that these dimers were indeed formed by the specific disulfide linkage between C-terminal cysteines.

Example 5: Animal Studies

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Leukemia is likely to be successfully treated using radiolabled anti-CD19 scFv because it is radiosensitive and there is ready access of antibody to the marrow space. Clinical studies have shown that iodine-labled antiferritin antibodies provided symptomatic relief to 77% patients with refractory Hodgkin disease and produced objective tumor regression in 40% of patients. In another clinical trail, when radiolabled anti-CD33 and -35 antibodies were used in combination with high a dose of cyclophosphamide, an overall of 19% complete remissions and 75% partial remissions were achieved for 210 evaluable patients with hematologic malignancies. The major side effect associated with the use of iodine-labled antibodies was reported to be thrombocytopenia, which occurred more frequently when the dose of iodine used was greater than 200 mCi/patient (see review by Grossbard *et al.*, 1992).

Radiolabled antibodies kill target cells by by-stander effect. Internalization of radiolabled antibodies is probably not desirable. It has shown that internalized radiolabled antibodies had a much shorter retention time and a faster rate of deiodination, which would dramatically reduce the efficacy of the therapeutic values of the antibodies (Richard et al., 1992). The single chain antibodies have the advantages of being small, with relatively high affinity toward the antigens and not being internalized by the target cells.

A. Preparation of FVS 191 and FVS 192 single chain antibody

The antibody is expressed in *Escherichia coli* as inclusion bodies. The inclusion bodes are denatured, refolded, and purified by FPLC chromatography. Since endotoxin contents of the antibody is high, it must be removed before being used in animals. Endotoxin is removed by affinity chromatography (a kit is commercially available). The amount of endotoxin in the antibody preparation is monitored by the Limulus Amebocyte Lysate Assay (Biowhittater Inc., Walkersville, MD). According to the US standard, the endotoxin contents in the final antibody preparation must be reduced to <15 endotoxin unit (EU, 1 EU = 0.5 ng/ml).

B. Iodination

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The single chain antibody is labeled with Na ¹³¹I using a Idogen kit (Pierce, Rockford, IL). The ratio of Idogen to antibody is adjusted to approximately 100ug:1mg as described by Badger et al. (1985). The labeled antibody will be separated from free ¹³¹I by gel filtration. The labeling efficiency and specific activity will be determined by cyclic anhydride method (Hantowich *et al.*, 1983). A specific activity of 1.0Ci/g or less should be suitable for the experiments. The same amounts of whole monoclonal antibody and Fab of an unrelated antibody should be labeled with ¹³¹I the same way to serve as controls.

C. Determination of immunoreactivity.

Immunoreactivity is defined as percentage of counts that are able to bind at antigen excess. Briefly, a serial dilution of target cells (CD19+, 106-7/ml will be incubated with labeled antibody (4-5 ng/ml) for 1 h at RT. Cells are centrifuged and supernatant radioactivity is counted. Immunoreactivity will be determined by Lineweaver-Burk analysis. Avidity of the antibody will be determined by incubation fixed amounts of

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cells (105/ml) with a serial dilutions of labeled antibody for 1 h at RT. Cells are washed and the cell pellet radioactivity is used to calculate the avidity (association constant and the number of binding site per cell).

D. In vitro measurement of single chain Fv metabolism.

This experiment determines the rate at which the labeled antibody is internalized and degraded. The target cells are incubated with labeled antibodies (scFv and whole Mab, 5 ng/106 cells) in a volume of 100 ul for 45 min on ice. The cells are washed and cultured at 37C. Aliquots of the incubation mixture are assayed for cell associated and supernatant radioactivity at 0, 1, 4, 10 and 24 h. the percentage of TCA precipitated radioactivity will be used for calculating the rate of internalization and intracellular metabolism of the labeled antibody.

1. Pharmacokinetic Studies

Pharmacokinetic studies are carried out by injecting labeled single chain antibody into a group of 4 BALB/c mice via the tail vein. Blood samples are collected at various time intervals. Radioactivities associated with the blood samples will be determined and T alpha 1/2 and T beta 1/2 of the single chain antibody will be calculated by computer simulation. As a control, the parental monoclonal antibody is labeled and injected into the mice as described above. Biodistribution is performed with paired labeling, e.g. the single chain antibody will be labeled with 131I and the controlled antibody labeled with ¹²⁵I. In our laboratory, anti CD19 scFv, FVS 191, has been successfully labled with 125I and used in immunochemistry and pharmacokinetics studies. Using current protocal, this scFv can be readily labled with 125I with a specific activity of 2.4 mci/mg. The immunoreactivity of the labled antibody was 55%. FVS 191 is more resistant to labeling damage than intact antibody. Results of Scatchard analysis showed that the affinity of FVS 191 toward CD19 antigen was 7.2 X 108 M⁻¹. This value is about four fold higher than its parent monoclonal antibody (1.93 X 108 M-1), suggesting that scFv may be a better targeting reagent than intact antibody. The observation that scFv showed higher affinity than its parent intact monoclonal antibody is

consistent with the findings of others. Data from pharmacokinetic studies in BALB/c mice showed that FVS 191 had a $T_{1/2}A$ and $T_{1/2}B$ of 2.5 min and 3.7 h respectively. In comparison, the intact monoclonal antibody had a $T_{1/2}A$, and $T_{1/2}B = 7.2$ min. and 57.1 h. In summary, the high specific immunoreactivity, high affinity and the rapid blood clearance of anti FVS 191 makes it an excellent candidate for use in cancer therapy.

2. Biodistribution Studies

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A mixture of equivalent amounts of specific antibody and control antibody with varied concentrations is injected i.v. into a group of 4 mice with human leukemia xenografts. The animals are sacrificed at 1, 24 and 48 h after the injection. Samples of blood, tumor, lung, spleen and kidney are weighed and counted in a gamma counter. The percentage of injected dose per gram of tissue (%ID/g) for each isotope is calculated. For dose escalation studies a single labeling (131I) will be performed to determine the proper dose range for subsequent animal survival tests.

E. Demonstration of Therapeutic Efficacy

Two types of leukemia animal models are used in the experiments — *e.g.* acute human leukemia (B1 or RS4:11 cell) in SCID mice or human acute leukemia xenograft tumor model in SCID or athymic BALB/c mice. The human leukemia SCID model has been well established in this laboratory and should be readily available for the experiments. The xenograft tumor model is established by injecting human leukemia cells (4-5 X 10⁷ in 0.2 ml PBS) into flanks of the mice as described by Richard et al, 1992). A palpable tumor module of 0.5-1.0 cm should be detected 8-10 days after the tumor cell injection. A single infusion (i.v) of various concentrations (low, medium and high) of radiolabled antibody is given to a group of 4 animals. The same amount of controlled antibody labeled with ¹³¹I are treated the same way. The percentage of survival will be recorded up to 50 days. For animals with xenografts, regression of tumors will be recorded. The definition of complete, and partial regressions needs to be defined.

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1. Therapeutic results

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Since FVS 191 and FVS 192 single chain antibody are specific for CD19+ cells, radiolabled antibody should show significant target cell killing effect in comparison to control antibody. Complete or partial tumor regression after radiolabled antibody treatment is expected. Due to its small size, single chain antibody is expected to penetrate the tumor more efficiently and show better results as compared to labeled whole MAb.

2. Therapy of Human B Cell Cancer (leukemia and lymphoma)

The anti CD19 scFv will be conjugated to ¹³¹I as described in animal therapy studies. Initial human trials will focus on pharmacokinetic and biodistribution studies.

Anti-CD19 antibodies have been effective for the treatment of human B cell leukemias or lymphomas when conjugated to toxins, e.g., ricin or pokeweed antiviral protein (Vitetta, et al., Uckun, et al.). B cell antibodies other than CD19 (e.g., anti-CD29) have been effective when linked to radioisotopes, e.g. 131I. Experience to date indicates that anti-CD19 FVS 191 and FVS 192 are not internalized by the cell after binding and thus these scFvshould be effective radioimmunoconjugates which should remain on the cell surface for optimal stability and cell killing. The anti-CD19 scFv will have very efficient biodistribution and tissue penetration based on the small size and short half life. As noted earlier, FVS 191 has a $T_{1/2}$ of 2.5 minutes in the alpha phase and $T_{1/2}$ of 3.7 hour in the beta phase. Thus, the rapid clearance combined should allow excellent killing of essentially all B cell leukemias and lymphomas (99% of which bear CD19). The small size of the ¹³¹I scFv should allow excellent killing in marrow lymph nodes and extromedullary sites which often serve as sanctuaries for leukemia and lymphoma cells.

REFERENCES

The references listed below, and all references cited herein, are incorporated by reference to the extent that they supplement, explain, provide a background for, or teach methodology, techniques and/or

5 compositions employed herein.

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Brief Description of Sequences

5	The following list briefly identifies the sequences discussed in the specification and claims:													
	SEQ ID NO:1	5' Oligonucleotide used for PCR of heavy chain variable region												
10	SEQ ID NO:2	3' Oligonucleotide used for PCR of heavy chain variable region												
15	SEQ ID NO:3	3' Oligonucleotide used for PCR of heavy chain constant region												
15	SEQ ID NO:4	5' Oligonucleotide used for PCR of light chain variable region												
20	SEQ ID NO:5	3' Oligonucleotide used for PCR of light chain variable region												
	SEQ ID NO:6	3' Oligonucleotide used for PCR of light chain constant region												
25	SEQ ID NO:7	Linker DNA between variable light and heavy chain regions												
	SEQ ID NO:8	cDNA sequence of B43 Heavy chain												
30	SEQ ID NO:9	cDNA sequence of SJ25C1 Heavy chain												
	SEQ ID NO:10 cDNA sequence of BLY3 Heavy chain													

	SEQ ID NO:11	cDNA sequence of B43 Light chain
5	SEQ ID NO:12	cDNA sequence of SJ25C1 Light chain
3	SEQ ID NO:13	cDNA sequence of BLY3 Light chain
	SEQ ID NO:14	Protein sequence of B43 Heavy chain
10	SEQ ID NO:15	Protein sequence of SJ25C1 Heavy chain
	SEQ ID NO:16	Protein sequence of BLY3 Heavy chain
15	SEQ ID NO:17	Protein sequence of B43 Light chain
10	SEQ ID NO:18	Protein sequence of SJ25C1 Light chain
	SEQ ID NO:19	Protein sequence of BLY3 Light chain
20	SEQ ID NO:20	Protein sequence of single chain B43 antibody
	SEQ ID NO:21	Protein sequence of single chain SJ25C1 antibody
25	SEQ ID NO:22	Protein sequence of single chain BLY3 antibody
20	SEQ ID NO:23	cDNA sequence of single chain B43 antibody
	SEQ ID NO:24	cDNAsequence of single chain SJ25C1 antibody
30	SEQ ID NO:25	cDNA sequence of single chain BLY3 antibody
	SEQ ID NO:26	Protein sequence of modified single chain B43 antibody

	SEQ ID NO:27	Protein sequence of modified single chain SJ25C1 antibody
5	SEQ ID NO:28	Protein sequence of the dimer single chain B43 antibody
	SEQ ID NO:29	Protein sequence of the dimer single chain SJ25C1 antibody

SEQUENCE LISTING

(1) GENERAL INFORMATION:

5 (i) APPLICANT: Bejcek, Bruce E.

Wang, Duo

Uckun, Fatih M.

Kersey, John H.

10 (ii) TITLE OF INVENTION:

IMMUNOCONJUGATES FROM SINGLE-CHAIN VARIABLE REGION FRAGMENTS OF ANTI-CD19 ANTIBODIES

15 (iii) NUMBER OF SEQUENCES: 29

(iv) CORRESPONDENCE ADDRESS:

20 (A) ADDRESSEE: Patterson & Keough, P.A.

(B) STREET: 527 Marquette Avenue South, Suite 1200

(C) CITY: Minneapolis

25

35

(D) STATE: Minnesota

(E) COUNTRY: USA

30 (F) ZIP: 55455

(v) COMPUTER READABLE FORM:

(A) MEDIUM TYPE: Floppy diskette, 3.5 inch

(B) COUMPUTER: Apple Macintosh

(C) OPERATING SYSTEM: Apple Macintosh System 7.0

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			(D)	SOFTWARE:		WordPeri	ect	3.0	for	Macintosh
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			(B)	FILING DATE):					
10			(C)	CLASSIFICAT	'ION:					
		(viii)	ATTORNEY/AG	ENT INF	ORMATION	ī:			
15			(A)	NAME: Danie	l F. Co	ughlin,	Esq.			
			(B)	REGISTRATIO	N NUMBE	R: 36	,111			
			(C)	REFERENCE/D	OCKET N	UMBER:				
20		(ix)	TELEC	OMMUNICATION	INFORM	ATION:				
			(A)	TELEPHONE:	612/34	9-5759				
25			(B)	TELEFAX:	612/34	9-9266				
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			(D)	MODOLOGY.	1:5000					

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5			(B)	TITLE:	*		
			(C)	JOURNAL:	*		
10			(D)	VOLUME:	*		
10			(E)	ISSUE:	*		
			(F)	PAGES:	*		
15			(G)	DATE:	*		
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20			(I)	FILING DATE	S:	*	
			(J)	PUBLICATION	DATE:	*	
			(K)	RELEVANT RE	ESIDUES:	+	t
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- 60 -

			(D)	TOPOLOGY:		linear	
		(x)	PUBLI	CATION INFOR	MATION	:	
5			(A)	AUTHORS:	*		
			(B)	TITLE:	*		
10			(C)	JOURNAL:	*		
10			(D)	VOLUME:	*		
			(E)	ISSUE:	*		
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			(I)	FILING DATE	:	*	
			(J)	PUBLICATION	DATE:	*	
25			(K)	RELEVANT RE	SIDUES:	*	
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(C) STRANDEDNESS: single (D) TOPOLOGY: linear 5 (x) PUBLICATION INFORMATION: (A) AUTHORS: * 10 (B) TITLE: * (C) JOURNAL: * VOLUME: (D) 15 (E) ISSUE: (F) PAGES: 20 (G) DATE: (H) DOCUMENT NUMBER: * **(I)** FILING DATE: * 25 (J) PUBLICATION DATE: * (K) RELEVANT RESIDUES: * 30 (xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 3: AGGCTTACTA GTACAATCCC TGGGCACAAT 30 (5) INFORMATION FOR SEQ ID NO:4 35 (i) SEQUENCE CHARACTERISTICS:

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5			(C)	STRANDEDNES	S:	single	
3			(D)	TOPOLOGY:	linear		
			(x)	PUBLICATION	INFORM	ATION:	
10			(A)	AUTHORS:	*		
				TITLE:	*		
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(i)

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10			(C)	JOURNAL:	*	
			(D)	VOLUME:	*	
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			(K)	RELEVANT RE	SIDUES:	r
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			(C)	STRANDEDNES	SS:	singl	.e
10			(D)	TOPOLOGY:	linea	r	
		(x)	PUBL	CATION INFOR	MATION	:	
15			(A)	AUTHORS:	*		
			(B)	TITLE:	*	•	
			(C)	JOURNAL:	*		
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35			(K)	RELEVANT RE	SIDUES:	:	*
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10			(C)	STRANDEDNES	S:	double	e stranded
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- 66 -

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25	Val	Lys	Gln	Arg	Pro	Gly	Gln	Gly	Leu	Glu	Trp	Ile	Gly	Gln	Ile	Trp	
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	65					70					75					80	
35																	
			CGA														288
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5	GGG ACC ACG (GTC A	CC										351		
	Gly Thr Thr V														
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	(10) INFORMATION FOR SEQ ID NO:9														
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15	((A)	LENGTH	:	345										
	,	(B)	TVDF.		nucleic acid										
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20	((C)	STRANDEDNESS: double stranded												
20	((D)	TOPOLO	GY:	linea	r									
	(ii) M	MOLECU	JLE TYPI	Ξ:	cDNA										
25	((A)	DESCRI	PTION	:	Heav	y ch	ain	SJ25	C1 D	NA				
	(vii) I	MMEDI	ATE SOU	JRCE:											
	((A)	LIBRARY	<i>ไ</i> :	Anti (CD-19	hyb.	rido	mas						
30															
	((B)	CLONE:		SJ25C	l cel	1 li:	ne							
	(x)	PUBLICA	ATION	INFOR	MTIO:	N:								
35	(.	A)	AUTHORS	5:	*										

(B)

TITLE:

- 69 -

				(C)	J	OURN	AL:	*									
				(D)	V	OLUM	E:	*							•		
5				(E)	I	SSUE	:	*									
				(F)	P.	AGES	:	*									
10				(G)	D	ATE:		*									
10				(H)	D	OCUM:	ENT :	NUMB	ER:	*							
				(I)	F	ILIN	G DA'	TE:		*							
15				(J)	Pi	JBLI	CATI	ON D	ATE:	*							
				(K)	Rl	ELEV	NT I	RESII	DUES	:	*						
20	(x:	i) A	SEQU	JENCI	E DES	SCRII	PTIO	N: :	SEQ :	ID N	D: 9	•					
	CTC	GAG	TCT	GGG	GCT	GAG	CTG	GTG	AGG	ССТ	GGG	TCC	TCA	GTG	AAG	Aጥጥ	48
														Val			10
					5					10					15		
25	TCC	TGC	AAG	GCT	TCT	GGC	ТАТ	GCA	TTC	AGT	AGC	TAC	TGG	ATG	AAC	TGG	96
														Met			, ,
				20					25				_	30		<u> </u>	
	GTG	AAG	CAG	AGG	CCT	GGA	CAG	GGT	CTT	GAG	TGG	ATT	GGA	CAG	ATT	TAT	144
	Val	Lys	Gln	Arg	Pro	Gly	Gln	Gly	Leu	Glu	Trp	Ile	Gly	Gln	Ile	Tyr	
30			35					40					45				
	CCT	GGA	GAT	GGT	GAT	ACT	AAC	TAC	AAT	GGA	AAG	TTC	AAG	GGT	CAA	GCC	192
	Pro		Asp	Gly	Asp	Thr		Tyr	Asn	Gly	Lys		Lys	Gly	Gln	Ala	
35		50					55					60					
).)	ልሮክ	ריוזים	ል ር-ጥ	GCA	GAC	ΔΔΔ	ጥርር	ጥርር	AGC	ልሮአ	GCC	ጥልር	<u>ል</u> ጥር	CAG	CTTC	ACC	240
														Gln			240
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						100					-						

	GGC CTA ACA	TCT (GAG GAC	TCT	GCG	GTC	TAT	тст	TGT	GCA	AGA	AAG	ACC	288		
	Gly Leu Thr													200		
			35				90		_		J	95				
5																
	ATT AGT TO													336		
	Ile Ser Ser		/al Asp	Phe	Tyr		Asp	Asn	Trp	Gly		Gly	Thr			
		100				105					110					
10	ACG GTC ACC	ı.									345	5				
	.Thr Val Thr															
	115															
15	(11) INFOR	MATION ·	FOR SE	Q ID	NO:	10										
13	(i) SEQUENCE CHARACTERISTICS:															
	(1) DECORACE CHARACTERISTICS.															
		(A)	LENGTH	:	34	2										
•																
20		(B)	TYPE:		nu	nucleic acid										
		(C)	ss:	S: double stranded												
	(C) STRANDEDNESS: double stranded															
		(D)	TOPOLO	GY:	li	near										
25																
	(ii)	MOLEC	cD:	CDNA												
		(A)	DESCRI	ייים מר⊥ רייים	J.		uoar.	y ch	ain	כע זם	מזאר					
		(11)	DBBCRT	11101	••		neav.	y CII	ain	סנום	DIVA					
30	0 (vii) IMMEDIATE SOURCE:															
		(A)	LIBRAR	Υ:	An	ti C	D-19	hyb	rido	mas						
		(D)	CLONE:		יזם	v2 ~	011 ·	14~-								
35		(B)	CHONE:		מר.	io C	ETT .	TTUE								
		(x)	PUBLICA	MOITA	JINI	FORM	ATIO	N:								

(A)

AUTHORS:

- 71 -

(B) TITLE: (C) JOURNAL: .5 (D) VOLUME: (E) ISSUE: 10 (F) PAGES: (G) DATE: (H) DOCUMENT NUMBER: 15 (I) FILING DATE: (J) PUBLICATION DATE: * 20 RELEVANT RESIDUES: (K) (xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 10: CTC GAG TCT GGG GCT GAG CTG GTG AGG CCT GGG GCC TCA GTG AAG ATT 48 25 Leu Glu Ser Gly Ala Glu Leu Val Arg Pro Gly Ala Ser Val Lys Ile 5 10 15 TCC TGC AAA GCT TCT GGC TAC GCA TTC AGT AGC TCT TGG ATG AAC TGG 96 30 Ser Cys Lys Ala Ser Gly Tyr Ala Phe Ser Ser Ser Trp Met Asn Trp 20 25 30 GTG AAG CAG AGG CCT GGA CAG GGT CTT GAG TGG ATT GGA CGG ATT TAT 144 Val Lys Gln Arg Pro Gly Gln Gly Leu Glu Trp Ile Gly Arg Ile Tyr 35 35 40 45 CCT GGA GAT GGA GAT ACT AAC TAC AAT GGA AAG TTC AAG GAA GCG GCC 192

Pro Gly Asp Gly Asp Thr Asn Tyr Asn Gly Lys Phe Lys Glu Ala Ala

- 72 -

50 55 60 ACA CTG ACT GCA GAC AAA TCC TCC AGC ACA GCG TAC ATG CAG CTC AGC 240 Thr Leu Thr Ala Asp Lys Ser Ser Ser Thr Ala Tyr Met Gln Leu Ser 5 65 70 75 80 AGC CTG ACC TCT GTG GAC TCT GCG GTC TAT TCT TGT GCA ÁGA TCG GAG 288 Ser Leu Thr Ser Val Asp Ser Ala Val Tyr Ser Cys Ala Arg Ser Glu 85 90 95 10 TAT TGG GGT AAC TAC TGG GCT ATG GAC TAC TGG GGC CAA GGG ACC ACG 336 Tyr Trp Gly Asn Tyr Trp Ala Met Asp Tyr Trp Gly Gln Gly Thr Thr 100 105 110 15 GTC ACC 342 Val Thr (12) INFORMATION FOR SEQ ID NO:11 20 SEQUENCE CHARACTERISTICS: (i) (A) LENGTH: 342 (B) TYPE: nucleic acid 25 (C) STRANDEDNESS: single stranded (D) TOPOLOGY: linear 30 (ii) MOLECULE TYPE: CDNA (A) DESCRIPTION: Light chain B43 DNA (vii) IMMEDIATE SOURCE: 35

(A)

(B)

LIBRARY:

CLONE:

Anti CD-19 hybridomas

B43 cell line

				(x)	P	UBLI	CATI	ON I	NFOR	ITAM	ON:						
5				(A)	A	UTHO	RS:	*									
3				(B)	т	ITLE	:	*					•				
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				(J)	PI	UBLIC	CATIO	ON D	ATE:	*							
25				(K)	R	ELEV	ANT I	RESI	DUES	:	*						
.23	(xi)) A	SEQU	JENCI	E DES	SCRII	OITS	N: 5	SEQ :	ID NO	D: 13	l :					
	GAG	CTC	GTG	CTC	ACC	CAG	TCT	CCA	GCT	TCT	TTG	GCT	GTG	тст	СТА	GGG .	48
																Gly	
30					5					10					15	_	
	CAG	AGG	GCC	ACC	∆ידיכי	ጥርር	ጥርር	AAG	GCC	AGC	$C\Delta\Delta$	A C.Tr	CTT	CAT	ייים איים. ייים איים	C N M	96
			Ala														90
	011.	9	1114	20			Cyb	בעב	25	DCI	Q111	DCI	vai	30	тУт	ASD	
35														-			
	GGT	GAT	AGT	TAT	TTG	AAC	TGG	TAC	CAA	CAG	ATT	CCA	GGA	CAG	CCA	CCC	144
	Gly	Asp	Ser	Tyr	Leu	Asn	Trp	Tyr	Gln	Gln	Ile	Pro	Gly	Gln	Pro	Pro	
			35					40					45				

(vii) IMMEDIATE SOURCE:

- 74 -

PCT/US96/06941

	AAA	CTC	CTC	ATC	TAT	GAT	GCA	TCC	AAT	CTA	GTT	TCT	GGG	ATC	CCA	CCC	192
	Lys	Leu	Leu	Ile	Tyr	Asp	Ala	Ser	Asn	Leu	Val	Ser	Gly	Ile	Pro	Pro	
		50					55					60					
5																	
	AGG	TTT	AGT	GGC	AGT	GGG	TCT	GGG	ACA	GAC	TTC	ACC	CTC	AAC	ATC	CAT	240
	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Asn	Ile	His	
	65					70					75					80	
10	CCT	GTG	GAG	AAG	GTG	GAT	GCT	GCA	ACC	TAT	CAC	TGT	CAG	CAA	AGT	ACT	288
	Pro	Val	Glu	Lys	Val	Asp	Ala	Ala	Thr	Tyr	His	Cys	Gln	Gln	Ser	Thr	
					85					90					95		
	GAG	GAT	CCG	TGG	ACG	TTC	GGT	GGA	GGG	ACC	AAG	CTG	GAA	ATA	AAA	CGT	336
15	Glu	Asp	Pro	Trp	Thr	Phe	Gly	Gly	Gly	Thr	Lys	Leu	Glu	Ile	Lys	Arg	
				100					105					110			
	AGA	TCT															342
	Arg	Ser															
20																	
	(13)	1I	IFOR1	1ATIC	ON FO	OR SE	EQ II	NO:	:12								
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25		i)	_)	SEQU	JENCE	E CHA	ARACI	ERIS	STICS	5:							
25				<i>(</i> =)			_	2.5									
				(A)	T	ENGTH	1:	33	33								
				(D)	mz.	7 TO TO .					ھا۔						
				(B)	.1.7	PE:		m	ıclei	.c ac	eia						
30				(C)	cn	RANI	SELVIE	ecc.		doub	ole s	.+~~~	الممالية				
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		<i>(</i> i	.i)	MOLE	CULE	: ጥሃ፤	E.	CT.	ONA								
35		, 1	- /														
				(A)	DF	SCRI	PTIC	N:		Liah	ıt ch	ain	SJ25	С1 Г	NA		
				,													

(A) LIBRARY:

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Anti CD-19 hybridomas (B) CLONE: SJ25C1 cell line 5 (x)PUBLICATION INFORMATION: (A) AUTHORS: 10 (B) TITLE: (C) JOURNAL: (D) VOLUME: 15 (E) ISSUE: (F) PAGES: 20 (G) DATE: (H) DOCUMENT NUMBER: FILING DATE: (I) 25 (J) PUBLICATION DATE: * (K) RELEVANT RESIDUES: 30 (xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 12: GAG CTC GTG CTC ACC CAG TCT CCA AAA TTC ATG TCC ACA TCA GTA GGA 48 Glu Leu Val Leu Thr Gln Ser Pro Lys Phe Met Ser Thr Ser Val Gly 5 10 35 GAC AGG GTC AGC GTC ACC TGC AAG GCC AGT CAG AAT GTG GGT ACT AAT 96 Asp Arg Val Ser Val Thr Cys Lys Ala Ser Gln Asn Val Gly Thr Asn 20 25 30

	GTA	GCC	TGG	TAT	CAA	CAG	AAA	CCA	GGA	CAA	TCT	CCT	AAA	CCA	CTG	ATT	144
	Val	Ala	Trp	Tyr	Gln	Gln	Lys	Pro	Gly	Gln	Ser	Pro	Lys	Pro	Leu	Ile	
			35					40					45				
5																	
	TAC	TCG	GCA	ACC	TAC	CGG	AAC	AGT	GGA	GTC	CCT	GAC	CGC	TTC	ACA	GGC	192
	Tyr	Ser	Ala	Thr	Tyr	Arg	Asn	Ser	Gly	Val	Pro	Asp	Arg	Phe	Thr	Gly	
		50					55					60					
•																	
10	AGT	GGA	TCT	GGG	ACA	GAT	TTC	ACT	CTC	ACC	ATC	ACT	AAC	GTG	CAG	TCT	240
	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Thr	Ile	Thr	Asn	Val	Gln	Ser	
	65					70					75					80	
	AAA	GAC	TTG	GCA	GAC	TAT	TTC	TAT	TTC	TGT	CAA	TAT	AAC	AGG	TAT	CCG	288
15	Lys	Asp	Leu	Ala	Asp	Tyr	Phe	Tyr	Phe	Cys	Gln	Tyr	Asn	Arg	Tyr	Pro	
					85					90					95		
	TAC	ACG	TCC	GGA	GGG	GGG	ACC	AAG	CTG	GAA	ATA	AAA	CGT	AGA	TCT		333
	Tyr	Thr	Ser	Gly	Gly	Gly	Thr	Lys	Leu	Glu	Ile	Lys	Arg	Arg	Ser		
20				100					105					110			
	(14)	IN	IFORM	IATIC	ON FO	OR SE	EQ II	NO:	13								
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25		(i	.)	SEQU	JENCE	CHA	RACI	ERIS	TICS	: ,							
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				(D)	TС	POLO	GY ·	li	near	,							
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35		(i	i)	MOLE	CULE	TYP	E:	cD	NA								
			•														

(A)

DESCRIPTION:

Light chain BLY3 DNA

- 77 **-**

(vii) IMMEDIATE SOURCE:

(A) LIBRARY: Anti CD-19 hybridomas 5 (B) CLONE: BLY3 cell line (x)PUBLICATION INFORMATION: AUTHORS: (A) 10 (B) TITLE: JOURNAL: (C) 15 (D) VOLUME: (E) ISSUE: (F) PAGES: 20 (G) DATE: (H) DOCUMENT NUMBER: .25 (I) FILING DATE: (J) PUBLICATION DATE: * RELEVANT RESIDUES: (K) 30 (xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 13: GAG CTC GTG CTC ACC CAG TCT CCA GCT TCT TTG GCT GTG TCT CTA GGG 48 Glu Leu Val Leu Thr Gln Ser Pro Ala Ser Leu Ala Val Ser Leu Gly 35 5 10 15

- 78 -

	CAG	AGG	GCC	ACC	ATC	TCC	TGC	AGA	GCC	AGC	CAG	AGT	GTT	GAT	AAT	TAT	96
	Glņ	Arg	Ala	Thr	Ile	Ser	Cys	Arg	Ala	Ser	Gln	Ser	Val	Asp	Asn	Tyr	
				20					25					30			
5	GGC	ATT	AGT	TTT	ATG	AAÇ	TGG	TTC	CAA	CAG	AAA	CCA	GGA	CAG	CCA	CCC	144
	Gly	Ile	Ser	Phe	Met	Asn	Trp	Phe	Gln	Gln	Lys	Pro	Gly	Gln	Pro	Pro	
			35					40					45				
	AAA	CTC	CTC	ATC	TAT	GCT	GCA	TCC	AAC	CAA	GGA	TCC	GGG	GTC	CCT	GCC	192
10					Tyr												
	_	50			_		55				-	60	-				
	AGG	ጥጥጥ	AGT	GGC	AGT	GGG	тст	GGG	ACA	GAC	ጥጥር	AGC	СТС	AAC	ልጥር	СДТ	240
					Ser												240
15	65		501	013		70	501	017		1105	75	Der	Deu	11011	110	80	
10	05					, 0					, 5					80	
	CCT	ልጥር	GAG	GAG	GAT	CAT	አ ርጥ	GCA	አጥር	m v m	መመር	ምርጥ	CAC	C λ λ	7 CIII	7 7 C	200
					Asp												288
	110	Mec	GIU	GIU		дел	1111	AIG	Mec	_	FIIE	Cys	GIII	GIII		пур	
20					85					90					95		
20	a. a						~~~					~~~					
					ACG												336
	GIU	vai	Pro	_	Thr	Pne	GIY	GIĀ	_	Thr	гàг	Leu	GIu		Lys	Arg	
				100					105					110			
0.5																	
25																	
	AGA	TCT															342
	Arg	Ser															
	(15)	II	IFORI	IATIO	ON FO	OR SE	EQ II	NO:	14								
30																	
		(i	_)	SEQU	JENCE	E CHA	ARACI	ERIS	STICS	5:							
				(A)	LE	ENGTH	I:	11	.7								
35				(B)	TY	PE:		am	nino	acid	l						

(C)

STRANDEDNESS:

- 79 -

(D) TOPOLOGY: (ii) MOLECULE TYPE: protein 5 DESCRIPTION: Heavy chain B43 protein (A) (vii) IMMEDIATE SOURCE: (A) LIBRARY: Anti CD-19 hybridomas 10 (B) CLONE: B43 cell line (x)PUBLICATION INFORMATION: 15 (A) AUTHORS: * (B) TITLE: (C) JOURNAL: 20 (D) VOLUME: (E) ISSUE: 25 (F) PAGES: (G) DATE: (H) DOCUMENT NUMBER: * 30 (I) FILING DATE: * (J) PUBLICATION DATE: * 35 (K) RELEVANT RESIDUES:

(xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 14:

	Leu	Glu	Ser	Gly	Ala 5	Glu	Leu	Val	Arg	Pro 10	Gly	Ser	Ser	Val	Lys 15	Ile
5	Ser	Cys	Lys	Ala 20	Ser	Gly	Tyr	Ala	Phe 25	Ser	Ser	Tyr	Trp	Met 30	Asn	Trp
	Val	Lys	Gln 35	Arg	Pro	Gly	Gln	Gly 40	Leu	Glu	Trp	Ile	Gly 45	Gln	Ile	Trp
10	Pro	Gly 50	Asp	Gly	Asp	Thr	Asn 55	Tyr	Asn	Gly	Lys	Phe 60	Lys	Gly	Lys	Ala
15	Thr 65	Leu	Thr	Ala	Asp	Glu 70	Ser	Ser	Ser	Thr	Ala 75	Tyr	Met ·	Gln	Leu	Ser 80
	Ser	Leu	Arg	Ser	Glu 85	Asp	Ser	Ala	Val	Туr 90	Ser	Cys	Ala	Arg	Arg 95	Glu
20	Thr	Thr	Thr	Val 100	Gly	Arg	Tyr	Tyr	Tyr 105	Ala	Met	Asp	Tyr	Trp 110	Gly	Gln
	Gly	Thr	Thr 115	Val	Thr											
25	(16)	IN	IFORM	IATIC	N FC	R SE	Q II	NO:	15							
		(i	.)	SEQU	ENCE	СНА	RACT	ERIS	TICS	:						
30				(A)	LE	NGTH	:	11	.5							
				(B)	ТУ	PE:		am	ino	acid						
				(C)	ST	'RAND	EDNE	ss:		•						
35				(D)	TO	POLO	GY:									

protein

(ii) MOLECULE TYPE:

- 81 -

(A) DESCRIPTION: Heavy chain SJ25C1 protein (vii) IMMEDIATE SOURCE: 5 (A) LIBRARY: Anti CD-19 hybridomas (B) CLONE: SJ25C1 cell line (x) PUBLICATION INFORMATION: 10 AUTHORS: (A) (B) TITLE: 15 (C) JOURNAL: (D) VOLUME: (E) ISSUE: 20 (F) PAGES: (G) DATE: 25 (H) DOCUMENT NUMBER: (I) FILING DATE: PUBLICATION DATE: * (J) 30 (K) RELEVANT RESIDUES: (xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 15: Leu Glu Ser Gly Ala Glu Leu Val Arg Pro Gly Ser Ser Val Lys Ile 35 10 15 Ser Cys Lys Ala Ser Gly Tyr Ala Phe Ser Ser Tyr Trp Met Asn Trp 20 25

Val Lys Gln Arg Pro Gly Gln Gly Leu Glu Trp Ile Gly Gln Ile Tyr 35 40 45 5 Pro Gly Asp Gly Asp Thr Asn Tyr Asn Gly Lys Phe Lys Gly Gln Ala 55 60 Thr Leu Thr Ala Asp Lys Ser Ser Ser Thr Ala Tyr Met Gln Leu Ser 65 70 75 80 10 Gly Leu Thr Ser Glu Asp Ser Ala Val Tyr Ser Cys Ala Arg Lys Thr 85 90 95 Ile Ser Ser Val Val Asp Phe Tyr Phe Asp Asn Trp Gly Gln Gly Thr 15 100 105 Thr Val Thr 115 20 (17) INFORMATION FOR SEQ ID NO:16 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 114 25 (B) TYPE: amino acid (C) STRANDEDNESS: 30 (D) TOPOLOGY: (ii) MOLECULE TYPE: protein . (A) DESCRIPTION: Heavy chain BLY3 protein 35 (vii) IMMEDIATE SOURCE:

(A) LIBRARY: Anti CD-19 hybridomas

			(B)	C	CLONE	:	В	LY3	cell	lin	ıe				
5			(x)	F	UBLI	CATI	ON I	NFOR	ITAM	ON:					
J			(A)	A	OHTU	RS:	*					٠			
			(B)	T	'ITLE	:	*								
10			(C)	J	OURN	AL:	*								
	•		(D)	v	OLUM	E:	*								٠
15			(E)	. I	SSUE	:	*								
15			(F)	P	AGES	:	*								
			(G)	D	ATE:		*								
20			(H)	D	OCUM	ENT 1	NUMB:	ER:	*						
			(I)	F	ILIN	G DA'	TE:		*						
25			(J)	P	UBLI	CATIO	ON D	ATE:	*						
2 .0			(K)	R	ELEV	ANT I	RESII	OUES	:	*					
	(xi)	A SEQ	UENC:	E DE	SCRI	PTIO	N: :	SEQ :	ID N	O: 10	6:				
30	Leu Gl	u Ser	Gly	Ala 5	Glu	Leu	Val	Arg	Pro 10	Gly	Ala	Ser	Val	Lys 15	Il€
35	Ser Cy:	s Lys	Ala 20	Ser	Gly	Tyr	Ala	Phe 25	Ser	Ser	Ser	Trp	Met 30	Asn	Trp
- •	Val Ly:	s Gln 35	Arg	Pro	Gly	Gln	Gly	Leu	Glu	Trp	Ile	Gly 45	Arg	Ile	Туг

- 84 -

Pro Gly Asp Gly Asp Thr Asn Tyr Asn Gly Lys Phe Lys Glu Ala Ala 50 55 Thr Leu Thr Ala Asp Lys Ser Ser Ser Thr Ala Tyr Met Gln Leu Ser 5 65 70 75 80 Ser Leu Thr Ser Val Asp Ser Ala Val Tyr Ser Cys Ala Arg Ser Glu 85 90 10 Tyr Trp Gly Asn Tyr Trp Ala Met Asp Tyr Trp Gly Gln Gly Thr Thr 100 105 Val Thr 15 (18) INFORMATION FOR SEQ ID NO:17 (i) SEQUENCE CHARACTERISTICS: 20 (A) LENGTH: 114 (B) TYPE: amino acid (C) STRANDEDNESS: 25 (D) TOPOLOGY: (ii) MOLECULE TYPE: protein 30 (A) DESCRIPTION: Light chain B43 protein (vii) IMMEDIATE SOURCE: (A) LIBRARY: Anti CD-19 hybridomas 35 (B) CLONE: B43 cell line

(x) PUBLICATION INFORMATION:

				(A)	A	UTHO	RS:	*								
5				(B)	Т	'ITLE	l:	*								
3				(C)	J	OURN	AL:	*								
				(D)	V	OLUM	E:	*								
10				(E)	I	SSUE	:	*								
				(F)	P	AGES	:	*								
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				(K)	R	ELEV	ANT :	RESI:	DUES	:	*					
25	(xi)) A	SEQ	UENC	E DE	SCRI	PTIO	N:	SEQ :	ID N	O: 1'	7:				
	Glu	Leu	Val	Leu	Thr 5	Gln	Ser	Pro	Ala	Ser 10	Leu	Ala	Val	Ser	Leu 15	Gly
30	Gln	Arg	Ala	Thr 20	Ile	Ser	Cys	Lys	Ala 25	Ser	Gln	Ser	Val	Asp 30	Tyr	Asp
	Gly	Asp	Ser 35	Tyr	Leu	Asn	Trp	Tyr 40	Gln	Gln	Ile	Pro	Gly 45	Gln	Pro	Pro
35	Lys	Leu 50	Leu	Ile	Tyr	Asp	Ala 55	Ser	Asn	Leu	Val	Ser 60	Gly	Ile	Pro	Pro
	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Asn	Ile	His

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65 70 75 80 Pro Val Glu Lys Val Asp Ala Ala Thr Tyr His Cys Gln Gln Ser Thr 85 90 95 5 Glu Asp Pro Trp Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys Arg 100 105 110 Arg Ser 10 (19) INFORMATION FOR SEQ ID NO:18 (i) SEQUENCE CHARACTERISTICS: 15 (A) LENGTH: 111 (B) TYPE: amino acid 20 (C) STRANDEDNESS: (D) TOPOLOGY: (ii) MOLECULE TYPE: protein 25 (A) DESCRIPTION: Light chain SJ25C1 protein (vii) IMMEDIATE SOURCE: 30 (A) LIBRARY: Anti CD-19 hybridomas (B) CLONE: SJ25C1 cell line PUBLICATION INFORMATION: (x)35 (A) AUTHORS:

(B)

TITLE:

				(C)	J	OURN	AL:	*								
5				(D)	Λ	OLUMI	Ξ:	*								
J				(E)	I	SSUE	:	*								
				(F)	P	AGES	:	*								
10				(G)	Di	ATE:		*								
				(H)	D	OCUMI	ENT I	NUMBI	ER:	*						
15				(I)	F.	ILING	G DA	re:		*						
				(J)	Ρί	JBLIC	CATIO	ON DA	ATE:	*						
				(K)	RI	ELEVA	ANT I	RESII	OUES	:	*					
20	(xi) A	SEQ	JENCI	E DES	SCRII	PTIO	J: S	SEQ I	ID NO	D: 18	3:				
	Glu	Leu	Val	Leu	Thr 5	Gln	Ser	Pro	Lys	Phe 10	Met	Ser	Thr	Ser	Val 15	GlΣ
25	Asp	Arg	Val	Ser 20	Val	Thr	Cys	Lys	Ala 25	Ser	Gln	Asn	Val	Gly 30	Thr	Asr
30	Val	Ala	Trp 35	Tyr	Gln	Gln	Lys	Pro 40	Gly	Gln	Ser	Pro	Lys 45	Pro	Leu	Ile
	Tyr	Ser 50	Ala	Thr	Tyr	Arg	Asn 55	Ser	Gly	Val	Pro	Asp 60	Arg	Phe	Thr	GlΣ
35	Ser 65	Gly	Ser	Gly	Thr	Asp 70	Phe	Thr	Leu	Thr	Ile 75	Thr	Asn	Val	Gln	Ser
	Lys	Asp	Leu	Ala	Asp 85	Tyr	Phe	Ťyr	Phe	Cys 90	Gln	Tyr	Asn	Arg	Tyr 95	Pro

	Tyr Thr Ser	Gly 0	Gly Gly Thr L	ys Leu 105	Glu	Ile	Lys	Arg	Arg	Ser
5	(20) INFOR	MOITAM	FOR SEQ ID	NO:19						
	(i)	SEQUE	INCE CHARACTE	RISTICS	5:					
10		(A)	LENGTH:	114						
		(B)	TYPE:	amino	acid					
		(C)	STRANDEDNES	S:						
15		(D)	TOPOLOGY:							
	(ii)	MOLEC	ULE TYPE:	protei	.n					
20		(A)	DESCRIPTION	:	Light	c ch	ain	BLY3	pro	tein
	(vii)	IMMED	IATE SOURCE:							
		(A)	LIBRARY:	Anti C	D-19	hyb:	rido	mas		
25		(B)	CLONE:	BLY3 c	ell]	line				
		(x)	PUBLICATION	INFORM	MOITA	1:				
30		(A)	AUTHORS:	*						
		(B)	TITLE:	*	٠					
		(C)	JOURNAL:	*						
35		(D)	VOLUME:	*		٠				
		(E)	ISSUE:	*						

- 89 -

(F) PAGES: (G) DATE: 5. (H) DOCUMENT NUMBER: * (I) FILING DATE: (J) PUBLICATION DATE: * 10 (K) RELEVANT RESIDUES: (xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 19: 15 Glu Leu Val Leu Thr Gln Ser Pro Ala Ser Leu Ala Val Ser Leu Gly 10 15 Gln Arg Ala Thr Ile Ser Cys Arg Ala Ser Gln Ser Val Asp Asn Tyr 20 20 25 Gly Ile Ser Phe Met Asn Trp Phe Gln Gln Lys Pro Gly Gln Pro Pro 35 40 45 25 Lys Leu Leu Ile Tyr Ala Ala Ser Asn Gln Gly Ser Gly Val Pro Ala 50 55 Arg Phe Ser Gly Ser Gly Thr Asp Phe Ser Leu Asn Ile His 65 70 75 80 30 Pro Met Glu Glu Asp Asp Thr Ala Met Tyr Phe Cys Gln Gln Ser Lys 85 90 95 Glu Val Pro Arg Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys Arg 35 100 105 110

Arg Ser

	(21)	INFOR	MATION	FOR SEQ ID	NO:20
5		(i)	SEQUE	NCE CHARACTE	RISTICS:
3			(A)	LENGTH:	246
			(B)	TYPE:	amino acid
10			(C)	STRANDEDNES	S:
			(D)	TOPOLOGY:	
15		(ii)	MOLEC	ULE TYPE:	protein
			(A)	DESCRIPTION	: single chain B43 protein
		(vii)	IMMED	IATE SOURCE:	
20			(A)	LIBRARY:	Anti CD-19 hybridomas
			(B)	CLONE:	B43 cell line
25			(x)	PUBLICATION	INFORMATION:
			(A)	AUTHORS:	*
			(B)	TITLE:	*
30			(C)	JOURNAL:	*
-			(D)	VOLUME:	*
35			(E)	ISSUE:	*
			(F)	PAGES:	*
			(G)	DATE:	*

				(H)	D	OCUM:	ENT I	NUMB:	ER:	*						
5				(I)	F.	ILIN	G DA'	ΓE:		*						
J				(J)	ΡŪ	UBLIC	CATI	ON D	ATE:	*						
				(K)	RI	ELEV	ANT I	RESI	DUES	: .	*					
10	(xi) A	SEQ	JENC!	E DES	SCRII	PTIO	N: ;	SEQ :	ID N	D: 20	0:				
	Leu	Glu	Ser	Gly	Ala 5	Glu	Leu	Val	Arg	Pro 10	Gly	Ser	Ser	Val	Lys 15	Ile
15	Ser	Cys	Lys	Ala 20	Ser	Gly	Tyr	Ala	Phe 25	Ser	Ser	Tyr	Trp	Met 30	Asn	Trp
20	Val	Lys	Gln 35	Arg	Pro	Gly	Gln	Gly 40	Leu	Glu	Trp	Ile	Gly 45	Gln	Ile	Trp
	Pro	Gly 50	Asp	Gly	Asp	Thr	Asn 55	Tyr	Asn	Gly	Lys	Phe	Lys	Gly	Lys	Ala
25	Thr 65	Leu	Thr	Ala	Asp	Glu 70	Ser	Ser	Ser	Thr	Ala 75	Tyr	Met	Gln	Leu	Ser 80
30	Ser	Leu _.	Arg	Ser	Glu 85	Asp	Ser	Ala	Val	Tyr 90	Ser	Cys	Ala	Arg	Arg 95	Glu
	Thr	Thr	Thr	Val	Gly	Arg	Tyr	Tyr	Tyr 105	Ala	Met	Asp	Tyr	Trp 110	Gly	Gln
35	Gly	Thr	Thr 115	Val	Thr	Gly	Gly	Gly 120	Gly	Ser	Gly	Gly ·	Gly 125	Gly	Ser	Gly
	Gly	Gly 130	Gly	Ser	Glu	Leu	Val	Leu	Thr	Gln	Ser	Pro	Ala	Ser	Leu	Ala

		Ser	Leu	Gly	Gln		Ala	Thr	Ile	Ser		Lys	Ala	Ser	Gln	
	145					150					155					160
5	Val	Asp	Tyr	Asp	Gly 165	Asp	Ser	Tyr	Leu	Asn 170	Trp	Tyr	Gln	Gln	Ile 175	Pro
	Gly	Gln	Pro	Pro 180	Lys	Leu	Leu	Ile	Tyr 185	Asp	Ala	Ser	Asn	Leu 190	Val	Ser
10																
	Gly	Ile	Pro 195	Pro	Arg	Phe	Ser	Gly 200	Ser	Gly	Ser	Gly	Thr 205	Asp	Phe	Thr
•	T. 211	Δen	Tlo	His	Pro	Val	Glu	Tare	77 ⇒ 1	λαρ	אן א	ת ת	mh w		II.	C
15	Dea	210	110	1115		Vai	215	БУЗ	Val	ASP	AIG	220	1111	īĀī	nis	Cys
	Gln	Gln	Ser	Thr	Glu	Asp	Pro	Trp	Thr	Phe	Gly	Gly	Gly	Thr	Lvs	Leu
	225					230					235		_		-	240
20	Glu	Ile	Lys	Arg	Arg	Ser										
					245											
25	(22)	IN	IFORM	IATIC	N FC	R SE	Q II	NO:	21							
		(i	١	SEOU	ENICE	СНА	ם א כית	י ד ס ידי	· ·ጥፕ C C							
		\ 1	. ,	2500	,	CIIA	irac i	LIVE	1103	•						
30				(A)	LE	NGTH	:	24	1							
				(B)	TY	PE:		am	ino	acid						
				(C)	ST	RAND	EDNE	SS:								
35				(D)	то	POLO	GY:									

(ii) MOLECULE TYPE: protein

- 93 -

(A) DESCRIPTION: single chain SJ25C1 protein (vii) IMMEDIATE SOURCE: 5 (A) LIBRARY: Anti CD-19 hybridomas (B) CLONE: SJ25C1 cell line PUBLICATION INFORMATION: (x)10 (A) AUTHORS: TITLE: (B) 15 (C) JOURNAL: (D) VOLUME: (E) ISSUE: 20 (F) PAGES: DATE: (G) 25 (H) DOCUMENT NUMBER: * (I) FILING DATE: PUBLICATION DATE: * (J) 30 (K) RELEVANT RESIDUES: (xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 21: 35 Leu Glu Ser Gly Ala Glu Leu Val Arg Pro Gly Ser Ser Val Lys Ile 5 10 15

	Ser	Cys	Lys	Ala	Ser	Gly	Tyr	Ala	Phe	Ser	Ser	Tyr	Trp	Met	Asn	${\tt Trp}$
				20					25					30		
5	Val	Lys	Gln 35	Arg	Pro	Gly	Gln	Gly 40	Leu	Glu	Trp	Ile	Gly 45	Gln	Ile	Tyr
10	Pro	Gly 50	Asp	Gly	Asp	Thr	Asn 55	Tyr	Asn	Gly	Lys	Phe 60	Lys	Gly	Gln	Ala
	Thr 65	Leu	Thr	Ala	Asp	Lys 70	Ser	Ser	Ser	Thr	Ala 75	Tyr	Met	Gln	Leu	Ser 80
15	Gly	Leu	Thr	Ser	Glu 85	Asp	Ser	Ala	Val	Tyr 90	Ser	Cys	Ala	Arg	Lys 95	Thr
	Ile	Ser	Ser	Val	Val	Asp	Phe	Tyr	Phe 105	Asp	Asn	Trp	Gly	Gln 110	Gly	Thr
20	Thr	Val	Thr 115	Gly	Gly	Gly	Gly	Ser 120	Gly	Gly	Gly	Gly	Ser 125	Gly	Gly	Gly
25	Gly	Ser 130	Glu	Leu	Val	Leu	Thr 135	Gln	Ser	Pro	Lys	Phe 140	Met	Ser	Thr	Ser
	Val 145	Gly	Asp	Arg	Val	Ser 150	Val	Thr	Cys	Lys	Ala 155	Ser	Gln	Asn	Val	Gly 160
30	Thr	Asn	Val	Ala	Trp 165	Tyr	Gln	Gln	Lys	Pro 170	Gly	Gln	Ser	Pro	Lys 175	Pro
	Leu	Ile	Tyr	Ser 180	Ala	Thr	Tyr	Arg	Asn 185	Ser	Gly	Val	Pro	Asp 190	Arg	Phe
35	Thr	Gly	Ser 195	Gly	Ser	Gly	Thr	Asp 200	Phe	Thr	Leu	Thr	Ile 205	Thr	Asn	Val

	Gln	Ser 210	Lys	Asp ·	Leu	Ala	Asp 215	Tyr	Phe	Tyr	Phe	Cys 220	Gln	Tyr	Asn	Arg
5	Tyr 225	Pro	Tyr	Thr	Ser	Gly 230	Gly	Gly	Thr	Lys	Leu 235	Glu	Ile	Lys	Arg	Arg 240
	Ser												,			
10	(23)	11	IFORI	OITAN	ON FO	OR SI	EQ II	ONO:	:22							
		(i	L)	SEQU	JENCI	E CH	ARACT	reris	STICS	S:						
15				(A)	LI ·	ENGTI	H:	24	13							
10				(B)	T	PE:		an	nino	ació	i					
				(C)	S	rani	DEDNE	ESS:								
20				(D)	TO	OPOLO	OGY:									
		(i	i)	MOLE	ECULI	E TYI	PE:	pı	rotei	in						
25 .				(A)	DI	ESCRI	IPTIC	ON:		sing	jle d	chair	ı BLY	73 pi	rotei	.n
23 .		(7)	/ii)	IMME	EDIAT	re so	OURCI	Ξ:								
				(A)	L	BRAI	RY:	Ar	nti C	CD-19) hyl	orido	omas			
30				(B)	CI	CONE	:	ві	LY3 c	cell	line	e				
				(x)	PŪ	JBLIC	CATIO	N IN	IFOR1	ATIC	N:					
35				(A)	ΙA	JTHOI	RS:	*								
<i></i>				(B)	T	TLE	:	*								
				(C)	JT,	NIRN	AL:	*								

				(D)	V	OLUM	E:	*								
5				(E)	I	SSUE	:	*								
	•		•	(F)	P	AGES	:	*								
				(G)	D	ATE:		*								
10				(H)	D	OCUM:	ENT :	NUMB	ER:	*						
				(I)	F	ILIN	g DA'	TE:		*						
15				(J)	P '	UBLI	CATI	ON D.	ATE:	*						
15				(K)	R	ELEV	ANT I	RESI	DUES	:	*					
	(xi) A	SEQ	JENCI	E DE.	SCRI	PTIO	N:	SEQ :	ID N	D: 2	2:				
20																
	Leu	Glu	Ser	Gly	Ala 5	Glu	Leu	Val	Arg	Pro 10	Gly	Ala	Ser	Val	Lys 15	Ile
25	Ser	Cys	Lys	Ala 20	Ser	Gly	Tyr	Ala	Phe 25	Ser	Ser	Ser	Trp	Met 30	Asn	Trp
	Val	Lys	Gln 35	Arg	Pro	Gly	Gln	Gly 40	Leu	Glu	Trp	Ile	Gly 45	Arg	Ile	Туг
30	Pro	Gly 50	Asp	Gly	Asp	Thr	Asn 55	Tyr	Asn	Gly	Lys	Phe 60	Lys	Glu	Ala	Ala
35	Thr 65	Leu	Thr	Ala	Asp	Lys 70	Ser	Ser	Ser	Thr	Ala 75	Tyr	Met	Gln	Leu	Ser
	Ser	Leu	Thr	Ser	Val 85	Asp	Ser	Ala	Val	Tyr 90	Ser	Cys	Ala	Arg	Ser 95	Glu

- 97 -

	Tyr	Trp	Gly	Asn 100	Tyr	Trp	Ala	Met	Asp 105	Tyr	Trp	Gly	Gln	Gly 110	Thr	Thr
5	Val	Thr	Gly 115	Gly	Gly	Gly	Ser	Gly 120	Gly	Gly	Gly	Ser	Gly 125	Gly	Gly	Gly
	Ser	Glu 130	Leu	Val	Leu	Thr	Gln 135	Ser	Pro	Ala	Ser	Leu 140	Ala	Val	Ser	Leu
10	Gly	Gln	Arg	Ala	Thr	Ile	Ser	Cys	Arg	Ala	Ser	Gln	Ser	Val	Asp	Asn
	145					150					155					160
15	Tyr	Gly	Ile	Ser	Phe 165	Met	Asn	Trp	Phe	Gln 170	Gln	Lys	Pro	Gly	Gln 175	Pro
	Pro	Lys	Leu	Leu 180	Ile	Tyr	Ala	Ala	Ser 185	Asn	Gln	Gly	Ser	Gly 190	Val	Pro
20	Ala	Arg	Phe 195	Ser	Gly	Ser	Gly	Ser 200	Gly	Thr	Asp	Phe	Ser 205	Leu	Asn	Ile
	His	Pro 210	Met	Glu	Glu	Asp	Asp 215	Thr	Ala	Met	Tyr	Phe 220	Cys	Gln	Gln	Ser
25	Lys 225	Glu	Val	Pro	Arg	Thr 230	Phe	Gly	Gly	Gly	Thr 235	Lys	Leu	Glu	Ile	Lys 240
	Arg	Arg	Ser													
30	(24)	II	IFORM	ATI(ON FO	OR SE	EQ II	ONO:	: 23							
		i)	L)	SEQU	JENCE	E CHA	ARACI	reris	STICS	S:						
35				(A)		ENGTH	H:	73								
				(B)	.1, 7	PE:		m	ıclei	.c ac	i⊥α					

- 98 -

		(C)	STRANDEDNES	S: double stranded
		(D)	TOPOLOGY:	linear
5	(ii)	MOLEC	ULE TYPE:	cDNA .
		(A)	DESCRIPTION	: single chain B43 DNA
10	(vii)	IMMED	IATE SOURCE:	
	·	(A)	LIBRARY:	Anti CD-19 hybridomas
		(B)		B43 cell line
15		(x)	PUBLICATION	INFORMATION:
		(A)	AUTHORS:	*
20		(B)	TITLE:	*
		(C)	JOURNAL:	*
		(D)	VOLUME:	*
25		(E)	ISSUE:	*
		(F)	PAGES:	*
30		(G)	DATE:	*
		(H)	DOCUMENT NUM	IBER: *
		(I)	FILING DATE:	*
35		(J)	PUBLICATION	DATE: *
		(K)	RELEVANT RES	IDUES: *

(x:	L) A	SEOUENCE	DESCRIPTION:	SEO	TD	$NO \cdot$	23.

	CTC	GAG	TCT	GGG	GCT	GAG	CTG	GTG	AGG	CCT	GGG	TCC	TCA	GTG	AAG	ATT	48
	Leu	Glu	Ser	Gly	Ala	Glu	Leu	Val	Arg	Pro	Gly	Ser	Ser	Val	Lys	Ile	
5					5					10					15		
	TCC	TGC	AAG	GCT	TCT	GGC	TAT	GCA	TTC	AGT	AGC	TAC	TGG	ATG	AAC	TGG	96
	Ser	Cys	Lys	Ala	Ser	Gly	Tyr	Ala	Phe	Ser	Ser	Tyr	Trp	Met	Asn	Trp	
		•		20					25					30			
10							٠										
	GTG	AAG	CAG	AGG	CCT	GGA	CAG	GGT	CTT	GAG	TGG	ATT	GGA	CAG	ATT	TGG	144
	Val	Lys	Gln	Arg	Pro	Gly	Gln	Gly	Leu	Glu	Trp	Ile	Gly	Gln	Ile	Trp	
			35					40					45				
15	CCT	GGA	GAT	GGT	GAT	ACT	AAC	TAC	TAA	GGA	AAG	TTC	AAG	GGT	AAA	GCC	192
	Pro	Gly	Asp	Gly	Asp	Thr	Asn	Tyr	Asn	Gly	Lys	Phe	Lys	${\tt Gly}$	Lys	Ala	
		50					55					60					
	ACT	CTG	ACT	GCA	GAC	GAA	TCC	TCC	AGC	ACA	GCC	TAC	ATG	CAA	CTC	AGC	240
20	Thr	Leu	Thr	Ala	Asp	Glu	Ser	Ser	Ser	Thr	Ala	Tyr	Met	Gln	Leu	Ser	
	65					70					75					80	
	AGC	CTA	CGA	TCT	GAG	GAC	TCT	GCG	GTC	TAT	TCT	TGT	GCA	AGA	CGG	GAG	288
	Ser	Leu	Arg	Ser	Glu	Asp	Ser	Ala	Val	Tyr	Ser	Cys	Ala	Arg	Arg	Glu	
25					85					90					95		
	ACT	ACG	ACG	GTA	GGC	CGT	TAT	TAC	TAT	GCT	ATG	GAC	TAC	TGG	GGC	CAA	336
	Thr	Thr	Thr	Val	Gly	Arg	Tyr	Tyr	Tyr	Ala	Met	Asp	Tyr	Trp	Gly	Gln	
				100					105					110			
30																	
	GGG	ACC	ACG	GTC	ACC	GGA	GGC	GGT	GGC	TCG	GGC	GGT	GGC	GGC	TCG	GGT	384
	Gly	Thr	Thr	Val	Thr	Gly	Gly	Gly	Gly	Ser	Gly	Gly	Gly	Gly	Ser	Gly	
			115					120					125				
35		GGC															432
	Gly	Gly	Gly	Ser	Glu	Leu	Val	Leu	Thr	Gln	Ser	Pro	Ala	Ser	Leu	Ala	
		130					135					140					

- 100 -

	GTG	TCT	CTA	GGG	CAG	AGG	GCC	ACC	ATC	TCC	TGC	AAG	GCC	AGC	CAA	AGT	480
	Val	Ser	Leu	Gly	Gln	Arg	Ala	Thr	Ile	Ser	Cys	Lys	Ala	Ser	Gln	Ser	
	145					150					155					160	
5	GTT	GAT	TAT	GAT	GGT	GAT	AGT	TAT	TTG	AAC	TGG	TAC	CAA	CAG	ATT	CCA	528
	Val	Asp	Tyr	Asp	Gly	Asp	Ser	Tyr	Leu	Asn	Trp	Tyr	Gln	Gln	Ile	Pro	
					165					170					175		
	GGA	CAG	CCA	ccc	AAA	CTC	CTC	ATC	TAT	GAT	GCA	TCC	AAT	CTA	GTT	TCT	576
10	Gly	Gln	Pro	Pro	Lys	Leu	Leu	Ile	Tyr	Asp	Ala	Ser	Asn	Leu	Val	Ser	
				180					185					190			
	GGG	ATC	CCA	ccc	AGG	TTT	AGT	GGC	AGT	GGG	TCT	GGG	ACA	GAC	TTC	ACC	624
	Gly	Ile	Pro	Pro	Arg	Phe	Ser	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	
15			195					200					205				
	CTC	AAC	ATC	CAT	CCT	GTG	GAG	AAG	GTG	GAT	GCT	GCA	ACC	TAT	CAC	TGT	672
	Leu	Asn	Ile	His	Pro	Val	Glu	Lys	Val	Asp	Ala	Ala	Thr	Tyr	His	Cys	
20		210					215					220					
	CAG	CAA	AGT	ACT	GAG	GAT	CCG	TGG	ACG	TTC	GGT	GGA	GGG	ACC	AAG	CTG	720
	Gln	Gln	Ser	Thr	Glu	Asp	Pro	Trp	Thr	Phe	Gly	Gly	Gly	Thr	Lys	Leu	
	225					230					235					240	
25	GAA	ATA	AAA	CGT	AGA	TCT											738
	Glu	Ile	Lys	Arg	Arg	Ser											
					245												
30	(25)	TN	IFORM	f A TT C	ON FO	R SF	O TE	NO:	2.4								
00	(23)		or Ord	11111)IV I C	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	.V	, 110.	44								
		(i	.)	SEQU	JENCE	CHA	RACT	ERIS	TICS	S:							
0.5				(A)	LE	NGTH	I:	74	1								
35				(B)	TY	PE:		nu	ıclei	c ac	id						

(C)

STRANDEDNESS:

double stranded

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(D) TOPOLOGY: linear (ii) MOLECULE TYPE: CDNA 5 (A) DESCRIPTION: single chain SJ25C1 DNA (vii) IMMEDIATE SOURCE: 10 (A) LIBRARY: Anti CD-19 hybridomas (B) CLONE: SJ25C1 cell line (x)PUBLICATION INFORMATION: 15 AUTHORS: (A) (B) TITLE: 20 (C) JOURNAL: (D) VOLUME: (E) ISSUE: 25 (F) PAGES: (G) DATE: 30 (H) DOCUMENT NUMBER: * (I) FILING DATE: (J) PUBLICATION DATE: * 35 (K) RELEVANT RESIDUES: (xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 24:

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	CTC	GAG	TCT	فافاف	GC.I.	GAG	CTG	GTG	AGG	CCT	GGG	TCC	TCA	GTG	AAG	ATT	48
	Leu	Glu	Ser	Gly	Ala 5	Glu	Leu	Val	Arg	Pro 10	Gly	Ser	Ser	Val	Lys 15	Ile	
5																	
	TCC	TGC	AAG	GCT	TCT	GGC	TAT	GCA	TTC	AGT	AGC	TAC	TGG	ATG	AAC	TGG	96
	Ser	Cys	Lys	Ala	Ser	Gly	Tyr	Ala	Phe	Ser	Ser	Tyr	Trp	Met	Asn	Trp	
				20					25					30			
10					CCT												144
	Val	Lys		Arg	Pro	Gly	Gln		Leu	Glu	Trp	Ile	_	Gln	Ile	Tyr	
			35					40					45				
	CCT	GGA	GAT	GGT	GAT	ACT	AAC	TAC	AAT	GGA	AAG	TTC	AAG	GGT	CAA	GCC	192
15					Asp												
		50					55					60					
		•															
	ACA	CTG	ACT	GCA	GAC	AAA	TCC	TCC	AGC	ACA	GCC	TAC	ATG	CAG	CTC	AGC	240
	Thr	Leu	Thr	Ala	Asp	Lys	Ser	Ser	Ser	Thr	Ala	Tyr	Met	Gln	Leu	Ser	
20	65					70					75					80	
	000	Cm ³	202	mam	GAG	CAC	mcm	ccc	СШС	m x m	mcm	mcm.	CON	202	7 7 C	7.00	200
					Glu												288
	Gry	Deu	1111	Der	85	mp	DCI	niu	Vul	90	DCI	Cys	nia	n. g	95		
25																	
	ATT	AGT	TCG	GTA	GTA	GAT	TTC	TAC	TTT	GAC	AAC	TGG	GGC	CAA	GGG	ACC	336
	Ile	Ser	Ser	Val	Val	Asp	Phe	Tyr	Phe	Asp	Asn	Trp	Gly	Gln	Gly	Thr	
				100					105					110			
20																	
30																GGC	384
•	Thr	Val	115	GIĀ	Gly	GΤΆ	GIY	120	GIĀ	GTĀ	GIĀ	GIÀ	125	GIÀ	GIÀ	GIĀ	
			113					120			•		123				
	003	mc c	<i>~</i> ~ ~	OFF	OTT C	ama.	7.00	C 2 C	more.	003	7. 7. 7.	тт~	N TO C	mcc.	707	шаг	422
35			_													TCA	432
	GTĀ	ser	GIU	ьеи	Val	neu	TIIT	GIII	Set	FTO	ηλρ	FIIG	met	ಶ್ವ	TIIT	ser	
		130					135					140					

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	GTA	GGA	GAC	AGG	GTC	AGC	GTC	ACC	TGC	AAG	GCC	AGT	CAG	AAT	GTG	GGT	528
	Val	G1y	Asp	Arg	Val	Ser	Val	Thr	Cys	Lys	Ala	Ser	Gln	Asn	Val	Gly	
	145					150					155					160	
						•											
5	ACT	AAT	GTA	GCC	TGG	TAT	CAA	CAG	AAA	CCA	GGA	CAA	TCT	CCT	AAA	CCA	576
	Thr	Asn	Val	Ala	Trp	Tyr	Gln	Gln	Lys	Pro	Gly	Gln	Ser	Pro	Lys	Pro	
,					165					170					175		
	CTG	ATT	TAC	TCG	GCA	ACC	TAC	CGG	AAC	AGT	GGA	GTC	CCT	GAC	CGC	TTC	624
10	Leu	Ile	Tyr	Ser	Ala	Thr	Tyr	Arg	Asn	Ser	Gly	Val	Pro	Asp	Arg	Phe	
				180					185					190			
	ACA	GGC	AGT	GGA	TCT	GGG	ACA	GAT	TTC	ACT	CTC	ACC	ATC	ACT	AAC	GTG	672
	Thr	Gly	Ser	Gly	Ser	Gly	Thr	Asp	Phe	Thr	Leu	Thr	Ile	Thr	Asn	Val	
15			195					200					205				
	CAG	TCT	AAA	GAC	TTG	GCA	GAC	TAT	TTC	TAT	TTC	TGT	CAA	TAT	AAC	AGG	720
	Gln	Ser	Lys	Asp	Leu	Ala	Asp	Tyr	Phe	Tyr	Phe	Cys	Gln	Tyr	Asn	Arg	
		210					215					220					
2 0																	
	TAT	CCG	TAC	ACG	TCC	GGA	GGG	GGG	ACC	AAG	CTG	GAA	ATA	AAA	CGT	AGA	738
	Tyr	Pro	Tyr	Thr	Ser	Gly	Gly	Gly	Thr	Lys	Leu	Glu	Ile	Lys	Arg	Arg	
	225					230					235					240	
25																	
	TCT																741
	Ser																٠
30	(26)	IN	1FORM	IATIC	ON FO	R SE	Q II	O NO:	25								
		(i	L)	SEQU	JENCE	E CHA	RACI	ERIS	TICS	3:	•						
				(A)	LE	INGTH	I:	72	9								
35																	
				(B)	ТУ	PE:		nu	ıclei	.c ac	id						
											_						
				(C)	SI	RANI	EDNE	ESS:		douk	ole s	trar	ıded				

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(D) TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA 5 DESCRIPTION: single chain BLY3 DNA (A) (vii) IMMEDIATE SOURCE: 10 (A) LIBRARY: Anti CD-19 hybridomas (B) CLONE: BLY3 cell line (x)PUBLICATION INFORMATION: 15 (A) AUTHORS: * (B) TITLE: 20 (C) JOURNAL: (D) VOLUME: (E) ISSUE: 25 PAGES: (F) DATE: (G) 30 (H) DOCUMENT NUMBER: * (I) FILING DATE: * (J) PUBLICATION DATE: * 35 (K) RELEVANT RESIDUES:

(xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 25:

	CTC	GAG	TCT	GGG	GCT	GAG	CTG	GTG	AGG	CCT	GGG	GCC	TCA	GTG	AAG	ATT	48
	Leu	Glu	Ser	Gly	Ala	Glu	Leu	Val	Arg	Pro	Gly	Ala	Ser	Val	Lys	Ile	
5					5					10					15		
	TCC	TGC	AAA	GCT	TCT	GGC	TAC	GCA	TTC	AGT	AGC	тст	TGG	ATG	AAC	TGG	96
											Ser						
10				20					25					30		•	
	GTG	AAG	CAG	AGG	CCT	GGA	CAG	GGT	CTT	GAG	TGG	ATT	GGA	CGG	ATT	TAT	144
	Val	Lys	Gln	Arg	Pro	Gly	Gln	Gly	Leu	Glu	Trp	Ile	Gly	Arg	Ile	Tyr	
			35					40					45				
15																	
	CCT	GGA	GAT	GGA	GAT	ACT	AAC	TAC	AAT	GGA	AAG	TTC	AAG	GAA	GCG	GCC	192
	Pro	Gly	Asp	Gly	Asp	Thr	Asn	Tyr	Asn	Gly	Lys	Phe	Lys	Glu	Ala	Ala	
		50					55					60					
20	ACA	CTG	ACT	GCA	GAC	AAA	TCC	TCC	AGC	ACA	GCG	TAC	ATG	CAG	CTC	AGC	240
	Thr	Leu	Thr	Ala	Asp	Lys	Ser	Ser	Ser	Thr	Ala	Tyr	Met	Gln	Leu	Ser	
	65					70					75					80	
	AGC	CTG	ACC	TCT	GTG	GAC	TCT	GCG	GTC	TAT	TCT	TGT	GCA	AGA	TCG	GAG	288
25	Ser	Leu	Thr	Ser	Val	Asp	Ser	Ala	Val	Tyr	Ser	Cys	Ala	Arg	Ser	Glu	
					85					90					95		
	TAT	TGG	GGT	AAC	TAC	TGG	GCT	ATG	GAC	TAC	TGG	GGC	CAA	GGG	ACC	ACG	336
	Tyr	Trp	Gly	Asn	Tyr	Trp	Ala	Met	Asp	Tyr	Trp	Gly	Gln	Gly	Thr	Thr	
30				100					105					110			
											•						
	GTC	ACC	GGA	GGC	GGT	GGC	TCG	GGC	GGT	GGC	GGC	TCG	GGT	GGC	GGC	GGA	384
	Val	Thr	Gly	Gly	Gly	Gly	Ser	Gly	Gly	Gly	Gly	Ser	Gly	Gly	Gly	Gly	
35			115					120					125				

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	TCC	GAG	CTC	GTG	CTC	ACC	CAG	TCT	CCA	GCT	TCT	TTG	GCT	GTG	TCT	CTA	432
	Ser	Glu	Leu	Val	Leu	Thr	Gln	Ser	Pro	Ala	Ser	Leu	Ala	Val	Ser	Leu	
		130					135					140					
5	GGG	CAG	AGG	GCC	ACC	ATC	TCC	TGC	AGA	GCC	AGC	CAG	AGT	GTT	GAT	ААТ	480
		Gln															
	145					150					155				-	160	
			•														
	TAT	GGC	ATT	AGT	TTT	ATG	AAC	TGG	TTC	CAA	CAG	AAA	CCA	GGA	CAG	CCA	528
10	Tyr	Gly	Ile	Ser	Phe	Met	Asn	Trp	Phe	Gln	Gln	Lys	Pro	Gly	Gln	Pro	
					165	,				170					175		
	CCC	AAA	CTC	CTC	ATC	TAT	GCT	GCA	TCC	AAC	CAA	GGA	TCC	GGG	GTC	CCT	576
	Pro	Lys	Leu	Leu	Ile	Tyr	Ala	Ala	Ser	Asn	Gln	Gly	Ser	Gly	Val	Pro	
15				180					185					190			
		AGG															624
	Ala	Arg		Ser	Gly	Ser	Gly		Gly	Thr	Asp	Phe	Ser	Leu	Asn	Ile	
20			195					200					205				
20																	
		CCT															672
	HIS	Pro	Met	GIU	GIU	Asp	_	unr	Ala	Met	Tyr		Cys	GIn	GIn	Ser	
		210					215					220					
25	AAG	CAC	CTIT	CCT	CCC	N C C	መመር	CCT	CCI	ccc	A CC	አአሮ	СПС	C A A	אַתעא	7 7 7	720
20		Glu															720
	225	Giu	vai	110	7 T G	230	1110	C±y	CLy	CLY	235	Lys	Deu	GIU	116	240	
	223					200					200					240	
	CGT	AGA	TCT														729
30	Arg																
	(27)	IN	FORM	IATIC	N FC	R SE	Q II	NO:	26		•						
		(i	.)	SEQU	ENCE	CHA	RACI	ERIS	TICS	S:							
35																	
				(A)	LE	NGTH	(:	24	7								

amino acid

(B)

TYPE:

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(C) STRANDEDNESS: (D) TOPOLOGY: 5 (ii) MOLECULE TYPE: protein (A) DESCRIPTION: modified single chain B43 antibody . 10 (vii) IMMEDIATE SOURCE: (A) LIBRARY: Anti CD-19 hybridomas 15 (B) CLONE: B43 cell line (x)PUBLICATION INFORMATION: (A) AUTHORS: * 20 (B) TITLE: (C) JOURNAL: **2**5 . (D) VOLUME: ISSUE: (E) PAGES: (F) 30 (G) DATE: (H) DOCUMENT NUMBER: * 35 (I) FILING DATE: * (J) PUBLICATION DATE: *

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(K) RELEVANT R	ESIDUES
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(xi)	Α	SEQUENCE	DESCRIPTION:	SEQ	ID	NO:	26:
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	,	,									· .	٠.				
5																
	Leu	Glu	Ser	Gly	Ala 5	Glu	Leu	Val	Arg	Pro	Gly	Ser	Ser	Val	Lys 15	Ile
10	Şer	Cys	Lys	Ala 20	Ser	Gly	Tyr	Ala	Phe 25	Ser	Ser	Tyr	Trp	Met 30	Asn	Trp
	Val		Gln 35	Arg	Pro	Gly	Gln	Gly 40	Leu	Glu	Trp	Ile	Gly 45	Gln	Ile	Trp
15	Pro	Gly 50	Asp	Gly	Asp	Thr	Asn 55	Tyr	Asn	Gly	Lys	Phe 60	Lys	Gly	Lys	Ala
20	Thr 65	Leu	Thr	Ala	Asp	Glu 70	Ser	Ser	Ser	Thr	Ala 75	Tyr	Met	Gln	Leu	Ser 80
	Ser	Leu	Arg	Ser	Glu 85	Asp	Ser	Ala	Val	Tyr 90	Ser	Cys	Ala	Arg	Arg 95	Glu
25	Thr	Thr	Thr	Val	Gly	Arg	Tyr	Tyr	Tyr 105	Ala	Met	Asp	Tyr	Trp 110	Gly	Gln
	Gly	Thr	Thr 115	Val	Thr	Gly	Gly	Gly 120	Gly	Ser	Gly	Gly	Gly 125	Gly	Ser	Gly
30	Gly	Gly 130	Gly	Ser	Glu	Leu	Val 135	Leu	Thr	Gln	Ser	Pro 140	Ala	Ser	Leu	Ala
35	Val 145	Ser	Leu	Gly	Gln	Arg 150	Ala	Thr	Ile	Ser	Cys 155	Lys	Ala	Ser	Gln	Ser 160
	Val	Asp	Tyr	Asp	Gly	Asp	Ser	Tyr	Leu	Asn	Trp	Tyr	Gln	Gln	Ile	Pro

170

175

165

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Gly Gln Pro Pro Lys Leu Leu Ile Tyr Asp Ala Ser Asn Leu Val Ser 180 185 190 Gly Ile Pro Pro Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr 5 195 . 200 205 Leu Asn Ile His Pro Val Glu Lys Val Asp Ala Ala Thr Tyr His Cys 210 215 10 Gln Gln Ser Thr Glu Asp Pro Trp Thr Phe Gly Gly Gly Thr Lys Leu 225 230 240 Glu Ile Lys Arg Arg Ser Cys 245 15 (28) INFORMATION FOR SEQ ID NO:27 (i) SEQUENCE CHARACTERISTICS: 20 (A) LENGTH: 242 (B) TYPE: amino acid (C) STRANDEDNESS: 25 (D) TOPOLOGY: (ii) MOLECULE TYPE: protein 30 (A) DESCRIPTION: modified single chain SJ25C1 antibody (vii) IMMEDIATE SOURCE: 35 (A) LIBRARY: Anti CD-19 hybridomas (B) CLONE: SJ25C1 cell line

- 110 -

				(x)	F	UBLI	CATI	ON :	INFO	RMATI	ON:					
				(A)	A	UTHO	DRS:	. ,	*							
5				(B)	Т	TTLE	Ξ:	1	*							
				(C)	J	OURN	IAL:	,	ŧ							
10				(D)	V	OLUM	Œ:	4	r							
				(E)	I	SSUE	:	,	•							
				(F)	P	AGES	:	*	,							
15				(G)	D.	ATE:		*	,							
				(H)	D	OCUM	ENT	NUME	BER:	*						
20				(I)	F	ILIN	g da	TE:		*						
				(J)	P	UBLI	CATI	ON D	ATE:	*						
				(K)	R	ELEV.	ANT :	RESI	DUES	:	*					
25	(xi)) A	SEQ	UENC	E DE:	SCRI	PTIO	N:	SEQ	ID N	0: 2	7:				
	Leu	Glu	Ser	Gly	Ala 5	Glu	Leu	Val	Arg	Pro 10	Gly	Ser	Ser	Val	Lys 15	Ile
30	Ser	Cys	Lys	Ala 20	Ser	Gly	Tyr	Ala	Phe 25	Ser	Ser	Tyr	Trp	Met 30	Asn	Trp
35	Val	Lys	Gln 35	Arg	Pro	Gly	Gln	Gly 40	Leu	Glu	Trp	Ile	Gly 45	Gln	Ile	Tyr
50	Pro	Gly 50	Asp	Gly	Asp	Thr	Asn 55	Tyr	Asn	Gly	Lys	Phe	Lys	Gly	Gln	Ala

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	Thr 65	Leu	Thr	Ala	Asp	Lys 70	Ser	Ser	Ser	Thr	Ala 75	Tyr	Met	Gln	Leu	Ser 80
5	Gly	Leu	Thr	Ser	Glu 85	Asp	Ser	Ala	Val	Tyr 90	Ser	Cys	Ala	Arg	Lys 95	Thr
	Ile	Ser	Ser	Val 100	Val	Asp	Phe	Tyr	Phe 105	Asp	Asn	Trp	Gly	Gln 110	Gly	Thr
10	Thr	Val	Thr 115	Gly	Gly	Gly	Gly	Ser 120	Gly	Gly	Gly	Gly	Ser 125	Gly	Gly	Gly
15	Gly	Ser 130	Glu	Leu	Val	Leu	Thr 135	Gln	Ser	Pro	Lys	Phe 140	Met	Ser	Thr	Ser
	Val 145	Gly	Asp	Arg	Val	Ser 150	Val	Thr	Cys	Lys	Ala 155	Ser	Gln	Asn	Val	Gly 160
20	Thr	Asn	Val	Ala	Trp 165	Tyr	Gln	Gln	Lys	Pro 170	Gly	Gln	Ser	Pro	Lys 175	Pro
	Leu	Ile	Tyr	Ser 180	Ala	Thr	Tyr	Arg	Asn 185	Ser	Gly	Val	Pro	Asp 190	Arg	Phe
25	Thr	Gly	Ser 195	Gly	Ser	Gly	Thr	Asp 200	Phe	Thr	Leu	Thr	Ile 205	Thr	Asn	Val
30	Gln	Ser 210	Lys	Asp	Leu	Ala	Asp 215	Tyr	Phe	Tyr	Phe	Cys 220	Gln	Туŗ	Asn	Arg
	Tyr 225	Pro	Tyr	Thr	Ser	Gly 230	Gly	Gly	Thr	Lys	Leu 235	Glu	Ile	Lys	Arg	Arg 240
35	Ser	Cys		٠												

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	(i)	SEQUE	NCE CHARACTE	RISTICS:
		(A)	LENGTH:	494
5		(B)	TYPE:	amino acid
		(C)	STRANDEDNES	S:
10		(D)	TOPOLOGY:	
	(ii)	MOLEC	ULE TYPE:	protein
15		(A)	DESCRIPTION antibody	dimer of single chain B43
15	(vii)	IMMED:	IATE SOURCE:	
		(A)	LIBRARY:	Anti CD-19 hybridomas
20		(B)	CLONE:	B43 cell line
		(x)	PUBLICATION	INFORMATION:
25		(A)	AUTHORS:	*
		(B)	TITLE:	*
		(C)	JOURNAL:	*
30		(D)	VOLUME:	* Y
		(E)	ISSUÈ:	*
35		(F)	PAGES:	*
		(G)	DATE:	*
		(H)	DOCUMENT NUM	MBER: *

				(I)	F	ILIN	G DA'	re:		*						
_				(J)	Pi	JBLI	CATIO	ON DA	ATE:	*						
5				(K)	RI	ELEV	ANT I	RESII	OUES	:	*					
	(xi) A	SEQ	JENCI	E DES	SCRII	PTIOI	N: \$	SEQ I	ID N	D: 21	В:				
10	Leu	Glu	Ser	Gly	Ala 5	Glu	Leu	Val	Arg	Pro	Gly	Ser	Ser	Val	Lys 15	Ile
15	Ser	Cys	Lys	Ala 20	Ser	Gly	Tyr	Ala	Phe 25	Ser	Ser	Tyr	Trp	Met 30	Asn	Trp
13	Val	Lys	Gln 35	Arg	Pro	Gly	Gln	Gly 40	Leu	Glu	Trp	Ile	Gly 45	Gln	Ile	Trp
2 0	Pro	Gly 50	Asp	Gly	Asp	Thr	Asn 55	Tyr	Asn	Gly	Lys	Phe 60	Lys	Gly	Lys	Ala
	Thr 65	Leu	Thr	Ala	Asp	Glu 70	Ser	Ser	Ser	Thr	Ala 75	Tyr	Met	Gln	Leu	Ser 80
2 5	Ser	Leu	Arg	Ser	Glu 85	Asp	Ser	Ala	Val	Tyr 90	Ser	Cys	Ala	Arg	Arg 95	Glu
30	Thr	Thr	Thr	Val 100	Gly	Arg	Tyr	Tyr	Туr 105		Met	Asp	Tyr	Trp 110	Gly	Gln
	Gly	Thr	Thr 115	Val	Thr	Gly	Gly	Gly 120	Gly	Ser	Gly	Gly	Gly 125	Gly	Ser	Gly
35	Gly	Gly 130	Gly	Ser	Glu	Leu	Val 135	Leu	Thr	Gln	Ser	Pro 140	Ala	Ser	Leu	Ala
	Val	Ser	Leu	Gly	Gln	Arg 150	Ala	Thr	Ile	Ser	Cys 155	Lys	Ala	Ser	Gln	Ser 160

	Val	Asp	Tyr	Asp	Gly 165	Asp	Ser	Tyr	Leu	Asn 170	Trp	Tyr	Gln	Gln	175	Pro
_																
5	Gly	Gln	Pro	Pro 180	Lys	Leu	Leu	Ile	Туr 185		Ala	Ser	Asn	Leu 190	Val	Ser
10	Gly	Ile	Pro 195	Pro	Arg	Phe	Ser	Gly 200	Ser	Gly	Ser	Gly	Thr 205	Asp	Phe	Thr
10	Leu	Asn 210	Ile	His	Pro	Val	Glu 215	Lys	Val	Asp	Ala	Ala 220	Thr	Tyr	His	Cys
15	Gln 225	Gln	Ser	Thr	Glu	Asp 230	Pro	Trp	Thr	Phe	Gly 235	Gly	Gly	Thr	Lys	Leu 240
	Glu	Ile	Lys	Arg	Arg 245	Ser	Cys	Cys	Ser	Arg 250	Arg	Lys	Ile	Glu	Leu 255	Lys
20	Thr	Gly	Gly	Gly 260	Phe	Thr	Trp	Pro	Asp 265	Glu	Thr	Ser	Gln	Gln 270	Cys	His
25	Tyr	Thr	Ala 275	Ala	Asp	Val	Lys	Glu 280	Val	Pro	His	Ile	Asn 285	Leu	Thr	Phe
	Asp	Thr 290	Gly	Ser	Gly	Ser	Gly 295	Ser	Phe	Arg	Pro	Pro 300	Ile	Gly	Ser	Val
30	Leu 305	Asn	Ser	Ala	Asp	Tyr 310	Ile	Leu	Leu	Lys	Pro 315	Pro	Gln	Gly	Pro	Ile 320
	Gln	Gln	Tyr	Trp	Asn 325	Leu	Tyr	Ser	Asp	Gly 330	Asp	Tyr	Asp	Val	Ser 335	Gln
35	Ser	Ala	Lys ·	Cys 340	Ser	Ile	Thr	Ala	Arg	Gln	Gly	Leu	Ser	Val 350	Ala	Leu
	Ser	Ala	Pro	Ser	Gln	Thr	Leu	Val	Leu	Glu	Ser	Gly	Gly	Gly	Gly	Ser

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Gly Gly Gly Gly Ser Gly Gly Gly Thr Val Thr Thr Gly Gln Gly Trp Tyr Asp Met Ala Tyr Tyr Tyr Arg Gly Val Thr Thr Thr Glu Arg Arg Ala Cys Ser Tyr Val Ala Ser Asp Glu Ser Arg Leu Ser Ser Leu Gln Met Tyr Ala Thr Ser Ser Ser Glu Asp Ala Thr Leu Thr Ala Lys Gly Lys Phe Lys Gly Asn Tyr Asn Thr Asp Gly Asp Gly Pro Trp Ile Gln Gly Ile Trp Glu Leu Gly Gln Gly Pro Arg Gln Lys Val Trp Asn Met Trp Tyr Ser Ser Phe Ala Tyr Gly Ser Ala Lys Cys Ser Ile Lys Val Ser Ser Gly Pro Arg Val Leu Glu Ala Gly Ser Glu Leu (30) INFORMATION FOR SEQ ID NO:29 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: (B) TYPE: amino acid (C) STRANDEDNESS:

(D)

TOPOLOGY:

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(ii) MOLECULE TYPE: protein DESCRIPTION: dimer of single chain SJ25C1 (A) 5 antibody (vii) IMMEDIATE SOURCE: LIBRARY: Anti CD-19 hybridomas (A) 10 (B) CLONE: SJ25C1 cell line (x)PUBLICATION INFORMATION: 15 **AUTHORS:** (A) (B) TITLE: (C) JOURNAL: 20 VOLUME: (D) ISSUE: (E) 25 (F) PAGES: (G) DATE: (H) DOCUMENT NUMBER: * 30 FILING DATE: (I) (J) PUBLICATION DATE: * 35 RELEVANT RESIDUES: (K) (xi) A SEQUENCE DESCRIPTION: SEQ ID NO: 29:

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	Leu	Glu	Ser	Gly	Ala 5	Glu	Leu	Val	Arg	Pro 10	Gly	Ser	Ser	Val	Lys 15	Ile
5	Ser	Cys	Lys	Ala 20	Ser	Gly	Tyr	Ala	Phe 25	Ser	Ser	Tyr	Trp	Met 30	Asn	Tr
٠	Val	Lys	Gln 35	Arg	Pro	Gly	Gln	Gly 40	Leu	Glu	Trp	Ile	Gly 45	Gln	Ile	Туз
10	Pro	Gly 50	Asp	Gly	Asp	Thr	Asn 55	Tyr	Asn	Gly	Lys	Phe 60	Lys	Gly	Gln	Ala
15	Thr 65	Leu	Thr	Ala	Asp	Lys 70	Ser	Ser	Ser	Thr	Ala 75	Tyr	Met	Gln	Leu	Ser
10	Gly	Leu	Thr	Ser	Glu 85	Asp	Ser	Ala	Val	Tyr 90	Ser	Cys	Ala	Arg	Lys 95	Thi
20	Ile	Ser	Ser	Val 100	Val	Asp	Phe	Tyr	Phe 105	Asp	Asn	Trp	Gly	Gln 110	Gly	Thr
	Thr	Val	Thr 115	Gly	Gly	Gly	Gly	Ser 120	Gly	Gly	Gly	Gly	Ser 125	Gly	Gly	GlΣ
25	Gly	Ser 130	Glu	Leu	Val	Leu	Thr 135	Gln	Ser	Pro	Lys	Phe 140	Met	Ser	Thr	Ser
30	Val 145	Gly	Asp	Arg	Val	Ser 150	Val	Thr	Cys	Lys	Ala 155	Ser	Gln	Asn	Val	Gl ₃
	Thr	Asn	Val	Ala	Trp 165	Tyr	Gln	Gln	Lys	Pro 170	Gly	Gln	Ser	Pro	Lys 175	Pro
35	Leu	Ile	Tyr	Ser 180	Ala	Thr	Tyr	Arg	Asn 185	Ser	Gly	Val	Pro	Asp 190	Arg	Phe
	Thr	Gly	Ser 195	Gly	Ser	Gly	Thr	Asp 200	Phe	Thr	Leu	Thr	Ile 205	Thr	Asn	Va]

	Gln			Asp	Leu	Ala			Phe	туг	Phe			Туг	: Asn	Arg
		210	,				215					220				
5	Tyr 225	Pro	Tyr	Thr	Ser	Gly 230		Gly	Thr	Lys	Leu 235		Ile	Lys	Arg	Arg 240
10	Ser	Cys	Cys	Ser	Arg 245	Arg	Lys	Ile	Glu	Leu 250		Thr	Gly	Gly	Gl _y 255	Ser
	Thr	Tyr	Pro	Tyr 260	Arg	Asn	Tyr	Gln	Cys 265	Phe	Tyr	Phe	Tyr	Asp 270		Leu
15	Asp	Lys	Ser 275	Gln	Val	Asn	Thr	Ile 280	Thr	Leu	Thr	Phe	Asp 285	Thr	Gly	Ser
	Gly	Ser 290	Gly	Thr	Phe	Arg	Asp 295	Pro	Val	Gly	Ser	Asn 300	Arg	Tyr	Thr	Ala
20	Ser 305	Tyr	Ile	Leu	Pro	Lys 310	Pro	Ser	Gln	Gly	Pro 315	Lys	Gln	Gln	Tyr	Trp 320
25	Ala	Val	Asn	Thr	Gly 325	Val	Asn	Gln	Ser	Ala 330	Lys	Cys	Thr	Val	Ser 335	Val
	Arg	Asp	Gly	Val 340	Ser	Thr	Ser	Met	Phe 345	Lys	Pro	Ser	Gln	Thr 350	Leu	Val
30	Leu	Glu	Ser 355	Gly	Gly	Gly	Gly	Ser 360	Gly	Gly	Gly	Gly	Ser 365	Gly	Gly	Gly
		Thr 370	Val	Thr	Thr	Gly	Gln 375	Gly	Trp	Asn	Asp	Phe 380	Tyr	Phe	Asp	Val
35	Val 385	Ser	Ser	Ile	Thr	Lys 390	Arg	Ala	Cys	Ser	Tyr 395	Val	Ala	Ser		Glu 400
	Ser	Thr	Leu	Glv	Ser	Leu	Gln	Met	Tvr	Ala	Thr	Ser	Ser	Ser	Lvs	Asn

WO 96/36360

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405 410 415

PCT/US96/06941

Ala Thr Leu Thr Ala Gln Gly Lys Phe Lys Gly Asn Tyr Asn Thr Asp
420 425 430

5

Gly Asp Gly Pro Tyr Ile Gln Gly Ile Trp Glu Leu Gly Gln Gly Pro 435 440 445

Arg Gln Lys Val Trp Asn Met Trp Tyr Ser Ser Phe Ala Tyr Gly Ser 10 450 455 460

Ala Lys Cys Ser Ile Lys Val Ser Ser Gly Pro Arg Val Leu Glu Ala 465 470 475 480

15 Gly Ser Glu Leu

:19 Protein sequence of BLY3 Light chain

What is claimed is:

- 1 1. An isolated and purified polynucleotide encoding a single chain
- 2 variable region polypeptide that binds to a CD19 antigen.
- 1 2. The isolated and purified polynucleotide of claim 1, wherein the
- 2 polypeptide encoded comprises an amino acid residue sequence according
- 3 to SEQ ID NO: 20, 21 or 22.
- 1 3. The isolated and purified polynucleotide of claim 1 wherein the
- 2 polynucleotide comprises a nucleotide sequence according to SEQ ID NO:
- 3 23, 24 or 25.
- 1 4. An isolated and purified polynucleotide comprising a nucleotide
- 2 base sequence that is identical or complimentary to a segment of at least 10
- 3 contiguous nucleotide bases of SEQ ID NO: 23, 24 or 25, wherein the
- 4 polynucleotide hybridizes to a polynucleotide that encodes a single chain
- 5 variable region polypeptide that binds to a CD19 antigen.
- 1 5. The isolated and purified polynucleotide of claim 4, wherein the
- 2 encoded polypeptide binds to a CD19 antigen with a K_a of at least 1 x 109
- $3 M^1$.
- 1 6. An isolated and purified polynucleotide comprising a nucleotide
- 2 base sequence that is identical or complimentary to a segment of at least
- 3 100 contiguous nucleotide bases of SEQ ID NO: 23, 24 or 25, wherein the
- 4 polynucleotide hybridizes to a polynucleotide that encodes a single chain
- 5 variable region polypeptide that binds to a CD19 antigen.
- 1 7. An isolated and purified single chain variable region polypeptide
- 2 that binds to a CD19 antigen.

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- 1 8. The isolated and purified polypeptide of claim 7, wherein the
- 2 polypeptide has a molecular weight of approximately 28 kDa.
- 1 9. The isolated and purified polypeptide of claim 7, wherein the
- 2 polypeptide binds to a CD19 antigen with a K_a of at least 1 x 109 M-1.
- 1 10. The isolated and purified polypeptide of claim 7, wherein the
- 2 polypeptide comprises an amino acid residue sequence according to SEQ
- 3 ID NO: 20, 21 or 22.
- 1 11. The isolated and purified polypeptide of claim 7, wherein the
- 2 polypeptide is further modified by the site specific insertion of a cysteine
- 3 residue at the C-terminus of the polypeptide.
- 1 12. A dimer of an isolated and purified single chain variable region
- 2 polypeptide, wherein the dimer is prepared by linking a first polypeptide of
- 3 claim 11 with a second polypeptide of claim 11 through a disulfide bond
- 4 between a C-terminus cysteine residue on each polypeptide.
- 1 13. An isolated and purified single chain variable region polypeptide
- 2 that binds to a CD19 antigen, wherein the polypeptide is prepared by a
- 3 process comprising the steps of:
- 4 (A.) cloning a DNA sequence that encodes the polypeptide
- 5 into an expression vector;
- 6 (B.) transforming *E. Coli* cells with the expression vector;
- 7 and
- 8 (C.) expressing the polypeptide in the transformed cells.
- 1 14. An immunoconjugate for the treatment of cancer comprising a
- 2 single chain variable region polypeptide that binds to a CD19 antigen,
- 3 wherein the polypeptide is linked to at least one cytotoxic agent.

- 1 15. The immunoconjugate of claim 14, wherein the polypeptide
- 2 comprises an amino acid residue sequence according to SEQ ID NO: 20, 21
- 3 or 22.
- 1 16. The immunoconjugate of claim 14, wherein the at least one
- 2 cytotoxic agent is selected from the group consisting of single chain, double
- 3 chain, and multiple chain toxins.
- 1 17. The immunoconjugate of claim 14, wherein the at least one
- 2 cytotoxic agent is a radionuclide selected from the group consisting of beta-
- 3 emitting metallic radionuclides, alpha emitters, and gamma emitters.
- 1 18. An immunoconjugate for the treatment of cancer comprising a
- 2 polypeptide of claim 12, wherein the polypeptide is linked to at least one
- 3 cytotoxic agent.
- 1 19. The immunoconjugate of claim 18, wherein the at least one
- 2 cytotoxic agent is selected from the group consisting of single chain, double
- 3 chain, and multiple chain toxins.
- 1 20. The immunoconjugate of claim 18, wherein the at least one
- 2 cytotoxic agent is a radionuclide selected from the group consisting of beta-
- 3 emitting metallic radionuclides, alpha emitters, and gamma emitters.
- 1 21. A process for preparing an immunoconjugate comprising a single
- 2 chain variable region polypeptide that binds to a CD19 antigen, wherein
- 3 the process comprises the steps of:
- 4 (D.) preparing the polypeptide according to a method comprising the steps of:
- i) cloning a DNA sequence that encodes the polypeptide into an expression vector;
- 8 ii) transforming E. coli cells with the expression

9	vector; and
10	iii) maintaining the transformed cells under
11	biological conditions sufficient for expression of the
12	polypeptide.
13	(E.) providing a suitable toxin; and
14	(F.) conjugating the polypeptide to the toxin.

- 1 22. The process of claim 21, wherein the process further comprises the
- 2 step of labelling the immunoconjugate with a radionuclide.
- 1 23. An immunoconjugate for the treatment of cancer comprising a
- 2 polypeptide of claim 12, wherein the polypeptide is linked to at least one
- 3 cytotoxic agent.
- 1 24. The immunoconjugate of claim 23, wherein the at least one
- 2 cytotoxic agent is selected from the group consisting of single chain, double
- 3 chain, and multiple chain toxins.
- 1 25. The immunoconjugate of claim 23, wherein the at least one
- 2 cytotoxic agent is a radionuclide selected from the group consisting of beta-
- 3 emitting metallic radionuclides, alpha emitters, and gamma emitters.
- 1 26. An immunoconjugate for the treatment of cancer comprising a
- 2 polypeptide of claim 12, wherein the polypeptide is linked to at least one
- 3 cytotoxic agent.
- 1 27. The immunoconjugate of claim 18, wherein the at least one
- 2 cytotoxic agent is selected from the group consisting of single chain, double
- 3 chain, and multiple chain toxins.
- 1 28. The immunoconjugate of claim 18, wherein the at least one
- 2 cytotoxic agent is a radionuclide selected from the group consisting of beta-

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- 3 emitting metallic radionuclides, alpha emitters, and gamma emitters.
- 1 29. A method for the treatment of cancer comprising the steps of:
- 2 (G.) selecting a patient evidencing symptoms of a B-cell 3 cancer, wherein the cancer is selected from the group consisting of 4 leukemia and B-cell lymphoma;
 - (H.) administering to the patient a therapeutically effective amount of the immunoconjugate of claim 22 in a biocompatible dosage form.

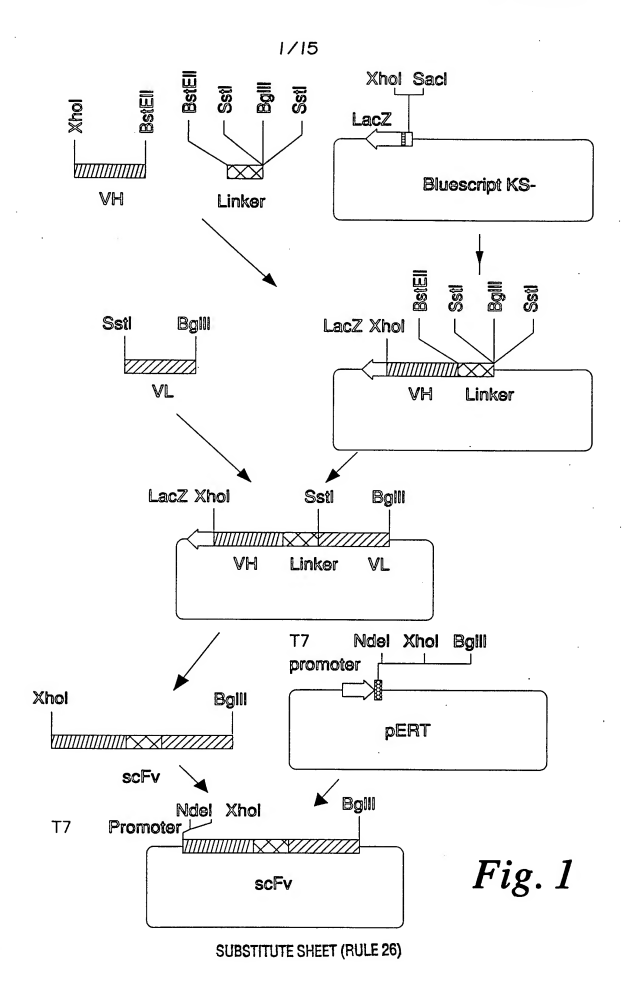


Fig. 2

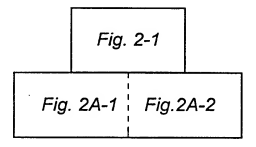
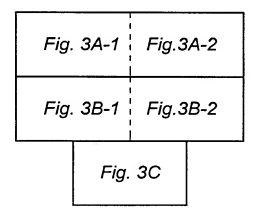


Fig. 3



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Heavy chain regions

CDR2 QIWPGDGDTNYNGKFKG Y R-YE	CDR2 DASNLVS S-TYRN- AQG-	
FR2 WVKQRPGQGLEWIG CDR3 KTISS-VDF-F S-YW GN W	ions FR2 DSYLN WYQQIPGQPPKLLIY AKSP I-FMFK	CDR3 QQSTED PWTFGG GTKLEIKRRS FCQYNRY-Y-S
**************************************	Light chain regions FR1 ELVLTQSPASLAVSLGQRATISC KASQSVDYDGDSYLNKFMST-V-G-V-VTN-GTNVA	FR3 GIPPRFSGSGSGTDFTLNIHPVEKVDAATYHC -V-DTT-TN-QSK-L-D-FY -V-ASM-ED-T-M-FC
B43 1 25C1 - BLY3 - B43 B43 BLY3 BLY3 B43 B43 BLY3 BLY3 BLY3 BLY3 A	B43 25C1 BLY3	B43 25C1 BLY3

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Fig. 24-1

CCT GGG TCC TCA GTG AAG ATT TCC TGC AAG GCT	G G	CDR2 AG TTC	CGGAT	ACT GCA GAC GAA TCC TCC AGC ACA GCC TAC ATG CAA CTC AGC AGC TTC TAT A G	A GG	ACG GTA GGC CGT TAT TAC TAT GCT ATG GAC TAC TGG TTA GATCTAC -AC	-GT AA-
FR1 CDR1 B43 CTC GAG TCT GGG GCT GAG CTG GTG AGG TCT GGC TAT GCA TTC AGT AGC TAC TGG ATG 25C1		FR2 B43 AAC TGG GTG AAG CAG AGG GGT GAT ACT AAC TAC AAT GGA A 25C1	BLY3	B43 AAG GGT AAA GCC ACT CTG CTA CGA TCT GAG GAC TCT GCG G 25C1 CA	-AA GCG T	FR3 CDR3 . B43 TCT TGT GCA AGA CGG GAG ACT ACG ACG GGC CAA GGG ACC ACG GTC ACC 25C1 AA- ACC -TGT T	BLY3 TC TA- TG-

Fig. 24-

	. CTC .	T CCT	
ν. - Ε	FR2 TGG TAC CAA CAG ATT CCA GGA CAG CCA CCC AAA CTC TAAA T-TTCA	C CAT	G GAA
-L	C AAA	C ATC	D
G AGG	A CCC	C AAC CC	2 4AG
G CAG	G CCA		A ACC
CTA GGG GA	A CAG	ACC	999
	GGA	1 DH 1 1	GGA
TCT	CCA	GACT	GGT A
GCT GTG T-C ACA	ATT -AA	-AA ACA	TTC -C-
	CAG	999 	ACG
TTG A	CAA	1 TCT 1 1	TGG -AC C
TCT 3AT -TC	TAC	-T- GGG	900 I I
CAG TCT CCA GCT TCT TTG GC CAA AGT GTT GAT AAA -TC A AG -G-	FR2 TGG	AGT	TAT
CCA AGT (TAT TTG AAC	T GGC TAT A	GAT AGG -T-
TCT CCA CAA AGT 	T TAT TTG AA(CTA GTT TCT	TTT A AGAC TTT AGT GGC GCA ACC TATC -CA GA GA	GAG A-C
CAG AGC C	 TAT TA G	TTT A AGA TTT AGT SCA ACC ' C -CA GA	ACT TA-
ບ ^ຂ , ,			AGT CAA
GAG CTC GTG CTC ACC TC TCC TGC AAG GCC 7	2 TAT GAT GGT GAT AGT TAT GAT GCA TCC AAT 0 1 AC- A TA -CC	1 9 1 4	et Et i
37. A	GGT GAT 3CA TCC 7	C AT GTG GATT GA- AAC TT GT G	CDR3 CAC TGT CAG CAA AAA CGT AGA TCT 1 TTA- TTC TGT 3 TT
CHC CC TC	3AT (T	GT (C)
FR1 GAG C TC TC 	-GA -GA T GA -CC A -	A T FR3 GGG ATC AG AAG (A G TCT AA	CAC TGT AAA CGT 7 TTA TT
AC I			B43 CAC ATA AAA 24C1 TT BLY3 TT-
B43 B43 ACC 25C1 A		HEET (RULE 26)	B43 ATA 24C BLY

CTG	FR2 TGG	AAT	CTA G G -	ATG T-T
GAG	AAC	TAC	AGC G	GCT TAC
GCT	ATG	AAC	AGC	TAT
9 1 1 1	TGG	ACT	CTC	TAC
TCT	TAC	GAT	CAA G	tAT
GAG	CDR1 AGC	GGT 	ATG 	cgt Gat t-g
CHC	AGT	GAT	TAC	Ggc -TA TAC
CTG	TTC 	GGA	000 000	GTA AAC
CAG	GCA	CCT	ACA	ACG t
949 1 1 1	TAT 	TGG -AT -AT	AGC	acg -gt tg'-
FR1 CAA	0 0 1 1	ATT 	TCC	act -t- ta-
A: B43 25C1 BLY3	B43 25C1 BLY3	B43 25C1 BLY3	B43 25C1 BLY3	B43 25C1 BLY3
	GTG CAG CTG CTC GAG TCT GGG GCT GAG	FR1 CAA GTG CTG GTG TCT GGG GCT GAG	FR1 CAA CTG CTC GAG TCT GGG GCT GAG GGC TAT GCA TTC AGC TAC TGG ATG AAC <td>FR1 CAA GTG CTG GAG TCT GGG GCT GAG </td>	FR1 CAA GTG CTG GAG TCT GGG GCT GAG

	TCT	CDR2 CAG	TCC	gag acc	
·	GCT	GGA	GAA A A	CDR3 cgg aa- tc-	
	AAG 	ATT 	GAC	AGA 	
	F	TGG	GCA : :	GCA 	
	10C	GAG 	ACT	TGT	ACC
	ATT 	CTT	CTG	H H H	0. H
•	AAG	GGT	ACT A A	TAT	ACG
Fig. 3A-2	GTG	CAG	0 0 1 1	GT - 1	A CC
ig. 3	TCA	GGA 	FR3 AAA C GCG	90 I I	9 9 1 1
F	TCC 9	CCT	GGT GAA	TCT :	CS
	999	AGG	AAG 	GAC 	00 i i
	CCT	CAG	11 TTC	GAG 	FR4
	AGG	AAG 	AAG	TC	TAC
	GTG 	GTG	GGA	CGA AC- ACC	GAC

	TTG A	TAT	FR3 GGG A	GAT C	ATA
	TCT -TC	AGT	TCT AG- C	GTG AAA -AT	GAA
	GCT AAA	GAT -CC AT-	GTT AAC -GA	AAG TCT G	CTG
	6 0	GGT -TA C	CTA -GG -A-	GAG C	AAG
<i>I-</i> 8	TCT	GAT A T	AAT T-C C	GTG 	ACC
Fig. 3B-1	CAG	TAT AC- A	TCC A	CCT	999
Fig	ACC	GAT G	GCA	CAT AC-	GGA G
	CTC	GTT G	CDR2 GAT TCG -C-	ATC	GGT A
	GTG	AGT -A-	TAT C	AAC -CC	FR4 TTC -C-
	CTC	CAA G	ATC T	CIC	ACG
	FR1 GAG	AGC T	CTC G	ACC T -G-	TGG AC C
	B: B43 25C1 BLY3	B43 25C1 BLY3	B43 25C1 BLY3	B43 25C1 BLY3	B43 25C1 BLY3

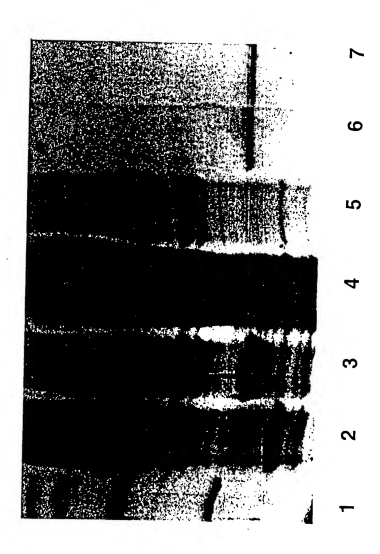
	0 0 1 1	CTC -CA	TTC	500 500	
	CDR1 AAG 	AAA 	GAC T	TAT	
	7GC	CCC	ACA	GAT AGG -T-	
	7CC 1 1 1 1 1 1 1 1 1	CCA T-T	9 9 9 9 1	CDR3 GAG A-C	
	ATC G	CAG A 	101 1 1 1	ACT TA- -AG	
2-2	ACC -G-	GGA	GGG A	AGT CAA 	
Fig. 3B-2	-L-	CCA	AGT	CAA TGT	
Fig	AGG	ATT -AA -AA	000	CAG TTC	
	CAG G-C	CAG	AGT -CA	TGT -A- 	
	GGG A	CAA	TTT C	CAC TT- TT-	
	CTA G	TAC T -T-	AGG C-C	TAT 	TCT
	TCT A	FR2 TGG	CCC GA- G	ACC GA- -TG	AGA
	GTG ACA	AAC	CCA T	GCA	CGT
	GC T	TTG	ATC G	GCT TTG A	AAA

Fig. 30

NY :	R-YE FR4 WGQGTTVT 	CDR2 PKLLIY DASNLVSP S-TYRN AQG-	GRS
FR2 WVKQRPGQGLEWIG 	 CDR3 RETTTVGRYYYAMDY KTISS-VDF-F S-YW -N- W	FR2 WYQQIPGQPPKLLIY KSP	FR4 FGGGTKLEIKRRS S
CDR1 SYWMN	CDR3 VYSCAR RETTT KTISS	CDR1 KASQSVDYDGDSYLN N-GTNVA RNY-I-FM-	CDR3 HC QQSTED PWT FY FCQYNRY-Y- FCK-V -R-
FR1 QVQLLESG AELVR	FR3 KATLTADESSSTAYMQLSSLRSEDSA QK	FR1 ELVLTQSP ASLAVSLGQRATISC KFMST-V-D-VSVT- 	FR3 GIPPRFSGSGSGTDFTLNIHPVEKVDAATYHC -V-DTT-TN-QSK-L-D-FY -V-ASM-ED-T-M-FC
A: B43 25C1	BLY3 B43 25C1 BLY3	B: B43 25C1 BLY3	B43 25C1 BLY3

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Fig. 4



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Fig. 5

Fig. 5A	Fig. 5B
Fig. 5C	Fig. 5D

13/15 **Fig. 5A**

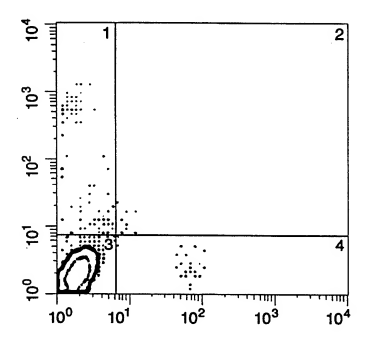
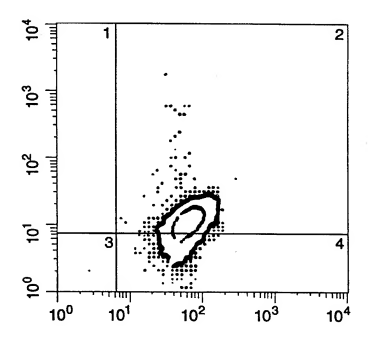


Fig. 5C



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Fig. 5B

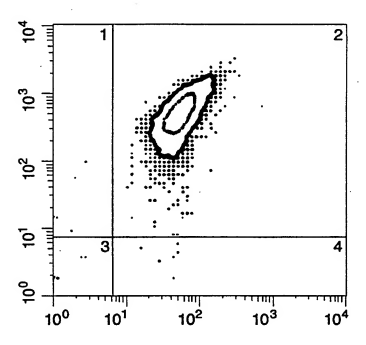
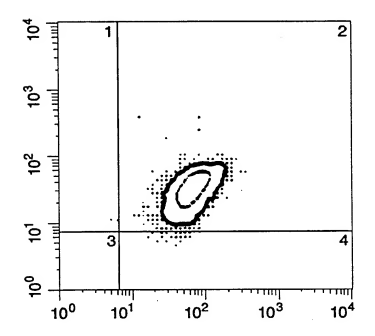
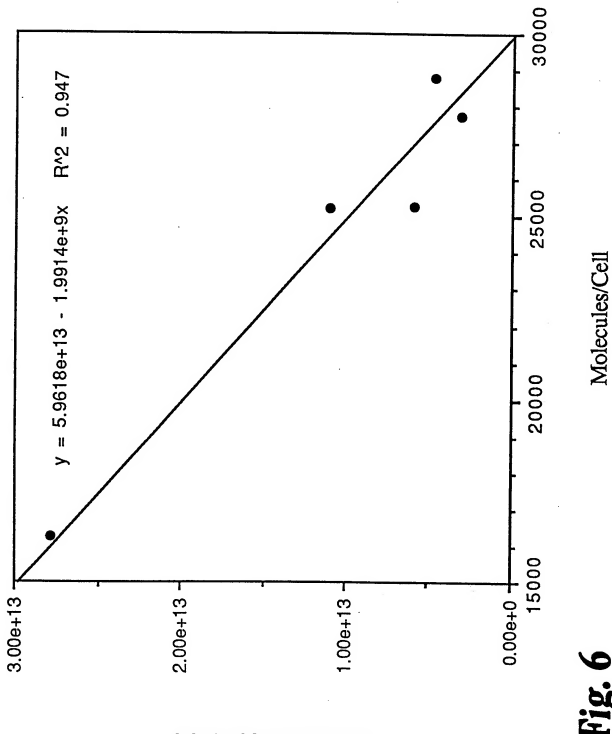


Fig. 5D



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Molecule L/Cell Mole

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INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/06941

IPC(6) US CL According t	SSIFICATION OF SUBJECT MATTER :A61K 39/395; C07K 16/42; C12N 1/20 :530/387.3, 388.22; 424/134.1 o International Patent Classification (IPC) or to both	national classification and IPC	
	ocumentation searched (classification system followed	by classification symbols)	
	530/387.3, 388.22; 424/134.1		XI.
Documentat	tion searched other than minimum documentation to the	e extent that such documents are included	in the fields searched
	lata base consulted during the international search (na llog, CAS, Sequence data bases	me of data base and, where practicable	, search terms used)
C. DOC	CUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.
Α	Science, Volume 267, No. 5199, Biotherapy of B-Cell Precursor Genestein to CD19-Associated Ty 11573419, see abstract.	Leukemia by Targeting	1-29
Υ	Cancer Reseach, Volume 51, No. Lambert et al., "An Immunotoxin Par A Natural Plant Toxin Adapted for 19057883, see abstract.	repared with Blocked Ricin	1-29
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X Furth	ner documents are listed in the continuation of Box C		
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International application No. PCT/US96/06941

C (Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where appropriate, of the releva	int passages	Relevant	to claim No
Y	Journal of Immunological Methods, Volume 136, No. 2 1991, Myers et al., "Roduction of Pokeweed Antiviral 1 PAP-containing immunotoxin B43-PAP directed against Human B Lineage Lymphoid Differentiation Antigen in Purified Form for Human Clinical Trials", abstract 817 abstract.	Protein the CD19 Highly	1-29	· · ·
A	Cancer Research, Volume 55, No. 11, issued 1995, Bej "Development and characterization of three recombinant chain antibody fragments (scFvs) directed against the CI antigen", abstract 11732023, see abstract.	dingle	1-29	
Ý.	US,A 5,091,513 (HUSTON ET AL.) 25 February 1992 document.	, see entire	1-29	
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